3

EUROPEAN PATENT APPLICATION published in accordance with Art. 158(3) EPC

EP 1 162 196 A1

(22) Date of filing: 22.12.2009 (43) Date of publication: 12.12.2001 Builetin 2001/50 (21) Application number: 00987728.3

(51) Int CL7: C07D 209/12, C07D 235/18, C07D 235/30, C07D 401/04, C07D 401/12, C07D 401/12, C07D 401/12, C07D 401/14, C07D 405/04, C07D 405/12, C07D 405/04, C07D 405/12, C07D 405/04, C07D 413/04, C07D 413/04, C07D 413/04, C07D 413/12, C07D 413/12, C07D 471/04, C07D 487/04

(86) International application number: PCT/JP00/09181

International publication number: WO 01/47883 (05.07.2001 Gazette 2001/27)

MIZUTANI, Kenji, Ctr. Pherm. Res. Inst. of Japan Tekatsuki-ehi, Oseka 569-1125 (JP) YOSHIDA, Atsuhito, Ctr. Pherm. Res. Inst. Japan

Takatsuki-ahi, Osaka 569-1125 (JP)

(84) Designated Contracting States:
AT BE CH CY DE DX ES FI FR GB GR IE IT LI LU
MC NL PT SE TR

AL LIT LY MK RO SI Designated Extension States:

(74) Representative: on Kreisler, Alek, Dipl.-Chem. et al

von Kreisier-Seiting-Werner Postfach 10 22 41 50482 Köln (DE)

Ê **FUSED-RING COMPOUNDS AND USE THEREOF AS DRUGS**

• HASHIMOTO, Hiromasa,

Ctr. Pharm. Res. Inst. Japan Takatsuku-ehi, Osaka 569-1125 (JP)

(71) Applicant: Jepan Tobacco Inc. Tokyo 105-8422 (JP) (30) Priority: 27.12.1999 JP 38900899

(57) The present invention provides a fused ring compound of the following formula [I]

a pharmaceutically acceptable sait hereol, and a ther-apeurite agent for hepatitis C, which conclain this com-pound. The compound of the present invention ahows an anti-hapatitis C virue (HCV) action besed on the HCV polymerase inhibitory activity, arm of is useful as a thera-pouric agent or prophylactic agent for hopatitis C. wherein each symbol is as defined in the specification,

Printed by Journ, 78001 PARIS (FR)

EP 1 162 198 A1

Technical Field

- (2001) The present invention relates to a novel fused ring compound and a pharmaceutically acceptable salt thereof useful da a therapeutic agent for hopatitis C. The present invention also rolates to a novel use of a certain fused ring compound or a pharmaceutically acceptable salt thereof as a therapeutic agent for hopatitis C. More particularly, the present invention relates to a therapeutic agent for hopatitis C, which contains a novel theat drig compound or a pharmaceutically acceptable salt thereof, which is effective for the prophylaxis or treatment of hepatitis C and which shows anti-hepatitia C virus (HCV) activity, particularly anti-HCV activity based on an RNA-depandent RNA polymerase
- Background Art

inhibitory activity.

õ

- (0002) In 1989, a main causalive virus of non-A non-B posttransfusion hepatitis was found and named hapatitis C virus (HCV). Since hen, several types of hopatitis viruses have been found boardes type A, type B and type C, wherein hapatitis caused by HCV is called hapatitis (1000). The patient of the world population, and the (0003) The patients infected with HCV are considered to involve several percent of the world population, and the
- infection with HCV characteristically becomes chronic.
- is oliminated by the immune system and the infection with this virus onds in an acute infection except for neonatos and infands having yet immetive immunological competence. In contrast, HCV somehow avoids the immune system of the host due to an unknown mechanism. Once infected with this virus, even an adult having a mature immune system [0004] HCV is an envelope RNA virus, whorein the genome is a single strand plus-strand RNA, and belongs to the genus Hepacivirus of Flavivirus (from The International Committee on Taxonomy of Viruses, International Union of Microbiological Societies). Of the same hapatits viruses, for exemple, hepatitis B virus (HBV), which is a DNA virus,
- frequently develops persistent infection. [0005] When chronic hepatits is associated with the persistent infection with HCV, it advances to cirrhesis or hepatic
- center in a high rate. Enucleation of tumor by operation does not help much, because the patient often develops recurrent hepatic center due to the sequels inflammation in non-centerorus parts.

 [0008] Thus, an effective therapeutic method of hepatitis C is desired. Apart from the symptomatic therapy to suppress inflammation with an anti-inflammatory agent, the development of a thorapeutic agent that reduces HCV to a low lovel free from inflammation and that oradicates HCV has been strongly demanded.

 [0007] At present, a treatment with freeferon is the only effective method known for the rest of the patient, it has no effect or provides the virus only in about one-third of the patient population, For the rest of the patients, it has no effect or provides only a temporary effect. Therefore, an anti-HCV drug to be used in the place of or concurrently with interferon is availed in great expectation.

 [0008] In record years, Ribevini (1-9-0-tholuraneyi-1+1-2-4-trazeto-2-texterorando) has become commercially in record years.
- scalable as attorapscute agent for hispetits C, which is to be used concurrently with interferon, it enhances the officery of interferon but only to a low efficery rate, and a different never thereportic agent of the patient. C is desired.

 [0099] Also, an attempt has been made to potentiate the formunocompetence of the patient with an interferon agentst, an interfewith-12 agents and the like, thereby to eradicate the virue, but an effective pharmaceuriscal agent has not
- been found yet.
- [0010] In addition, the inhibition of HCV growth, wherein HCV-specific protein is targeted, has been drawing attention
- (0011) The gene of HCV encodes a protein such as serine protease, RNA holicase, RNA depondent RNA polymentae and the like. These proteins function as a specific protein essential for the growth of HCV.

 [0012] One of the specific proteins, RNA-dependent RNA polymenses (hereinafter to be also briefly referred to as an HCV polymenses), is an enzyme essential for the growth of the virus. The gene replication of HCV having a plusaritand RNA specific protein start of RNA growth of the virus of the plusaritand RNA specific proteins and specific proteins and RNA specific proteins and sp like in other attempts to develop an anti-HCV drug based on other action mechanisms. As the situation stands,
- [0014] The following discloses known compounds rolatively similar to the compound of the present invention. [0015] A known therepeutio agent for hepatitis C having a benzimidazole skaleton is disclosed in WO97/36868

Jepanose Pstent Application under PCT laid-open under kohyo No. 2000-511898 (EP806097) and WO99/51619.
[0018] WO97/38888 discloses the following compound D and the like, and HCV helicase inhibitory activity of the

following compound E and the like, and WO98/51819 discloses the following compound F and the like, in both of which a possibility of these compounds being effective as an #CV inhibitor is mentioned.

[0018] However, these publications do not include the compound disclosed in the present specification, or a disclo-[0017] Japanese Patent Application under PCT taid-open under kohyo No. 2000-511898 (EP908097) discloses the

ō

[0019] A known anti-hepatitis virus agent having a benzimidazole skeleton is disclosed in Japanese Patent Applica-tion under PCT latd-open under kohye No. 2000-503017 (WOB725318) and Japanese Patent Application under PCT latd-open under kohye No. 10-505082 (WOB87848).

[0020] WC87/25316 discloses the following compound A and the like, wherein the use thereof is for a treatment of viral infection. The targot virus is a DNA virus such as hepatitis B virus and the like. However, this publication does not

include the compound disclosed in the present specification or a description regarding or suggestive of HCV. [0021] Japanese Patent Application under PCT laid-open under kohyo No. 10-505082 discloses the following comspecification or a description regarding or suggestive of HCV. as harpesvirus and hepatitis B virus. However, this publication does not include the compound disclosed in the present pound B and tho like, wherein the use thereof is for a treatment of viral infection. The target virus is a DNA virus such

Ġ

ŧ

[0023] The benzimidazole derivatives having an entiviral activity have been disclosed in JA-A-3-31264, US3044382 and US376504. In addition, WOS847672 discloses, as a production inhibitor of himor nocasis factor (TNF) and cyclic AMP, a benzimidazole derivative for the use as an anti-human immunodeficiency virus (HV) agont and an anti-inflammaniminodeficiency virus (HV) a

8

However, none of these publications includes the compound of the present invention or a description regarding

EP 1 162 198 A1

or suggestive of an anti-HCV effect.

(9024) Known benzinnidazole derivatives having a pharmacautized use other than as an antiviral agent are disclosed in JPA-46-601318 (USSS24651) and JP-A-9-104073 (USSS532429). These publications disclose the following compound C and the like as a catechol disther compound, and the use thereof as an anti-inflammation againt. However, notitiver of the publications includes the compound of the present invention, and as the action mechanism, the former discloses phosphodisterate by the did not include a description regarding or suggestive of an anti-HCV effect.

õ following compound G and the like, and the use thereof for the treatment of bronchitis, gromerutenephritis and the like. However, this publication does not include the compound of the present invention, but discloses only a phosphodiestenes of Vinhibitory and hypogycomic action. This publication does not include a description regarding or suggestive of [0025] Japanese Patent Application under PCT laid-open under kohyo No. 2000-159749 (EP882718) discipses the

ä

Ľ [0026] WO88/50026, WO68/50030 and WO68/50031 disclose benzimidazolo derivatives as an antitumor agent hav-ing a protein isopremyl transferase action. While this publication discloses a wide ecops of the claims, at least it does not include a compound analogous to the compound of the present invention or a description regarding or suggestive of an anti-HCV effect.

matory disease, and WO66/35713 discloses the application thereof as a growth hormone release promoter to treat a growth hormone-related disease such as esteoporests and the like. However, none of these publications includes a description regarding or suggestive of an anti-HCV effect.

[0028] JP-A-53-14735 discloses a benzimidazole derivative as a brightonor besides its pharmaceutical use, but this publication does not include the compound of the present invention. [0027] JP-A-8-109189 (EP894535) discloses the application of a tachykinin receptor antagonist to treat an inflam-

Disclosure of the invention

ĸ

(0029) Based on the findings from the proceding studies, it has been elucidated that a pharmaceutical agent having an anti-HCV earthy is effective for the prophysicis and treatment of hopatils. C, and particularly an enti-HCV agent having an inhibitory activity on RNA-dependent RNA pelymerase of HCV can be a prophysicis and therapoutic agent of a continuous properties. C and a prophysicio and therapoutic agent for the disease caused by hepatitis C. (0020) Accordingly, the present invention provides a pharmacoutical agent having an anti-HCV activity, particularly a pharmacoutical agent taking an RNA-dependent RNA polymerase inhibitory activity.

[0032] Thus, the present invention provides the following (1) to (43). RNA-dependent RNA polymeraso inhibitory activity, and completed the present invention.

à

å

(1) A therapsuite agent for hopathia C, which comprises a fused ring compound of the following formula [I] or a phermaceutically acceptable salt thereof as an active ingredient:

a broken line is a single bond or a double bond,

G1 G2 G3 G4 G5, G8, G8 and G9 G7 is C(-R*) or a nitrogen atom, are each independently a carbon atom or a nitrogen atom, are each independently a carbon atom or a nitrogen atom optionally substituted by R*, is C(-R*), an oxygen atom, a suitur atom, or a nitrogen atom optionally substituted by R*.

wherein R1, R2, R3 and R4 are each independently,

25

(1) hydrogen etom, (2) C₁₋₆ eikanoyl, (3) carboxyl,

(6) C₁₋₄ alkyl optionally substituted by 1 to 3 substituent(s) selected from the following group A. grup A: heliogen atom, hydroxyl group, carboxyl, amino, C₁₋₄ alkoxy, C₁₋₄ alkoxycarboxyl and C₁₋₆ alkylamino,

wherein R*1 is optionally substituted C_{I,8} alityf (as defined above) or C_{B,16} aryf C_{I,8} alityf optionally substituted by 1 to 5 autostituent(a) selected from the following group B, group B, group B, attorn, cyano, nitro, C_{I,8} alityf, alit

nalogenated C_{1,4} alkyl, C_{1,4} alkanoyl, (CH₂), NPP1PA2, (CH₂), NPP1-CORP2, (CH₂), NHSO₂RP1, (CH₂), ODP1, (CH₂), CONP1, (CH₂), SO₂RP1 and (CH₂), NPP1PA2, (CH₂), NPP1-CORP2, (CH₂), SO₂RP1 and (CH₂), SO₂NPP1PA2, (CH₂), SO₂RP1 and RP2 are each independently hydrogen atom or C_{1,4} alkyl and r is 0 or an integer of 1 to 8, (e) -CONPA2 RP2

mersin R42 and R43 are each independently hydrogen atom, $C_{1,6}$ alkoxy or optionally substituted $C_{1,6}$ alkyl

(as defined above), (9) -C(≃NR^{e4})NH₂

wherein Rad is hydrogen etom or hydroxyl group, (10) -NHRe5

Ġ

6

merein Ras is hydrogen atom, C14 alkanoyl or C14 alkylsullonyl,

wherein Re⁶ is hydrogen etem or optionally substituted C₁₋₈ alfy/(as dolined abova), (12) -SO₂,Re⁷ Theerein Re⁷ is hydroxyl group, amino, C₁₋₈ alkyl or C₁₋₈ alkylamino (11) -ORes

8

8

(13) -P(aO)(OR²⁶¹)₂ wherein R²⁶¹ is hydrogen atom, optionally substituted C₁₋₈ alkyl (as defined above) or C₆₋₁₄ aryl C₁₋₈ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, and

 \Re^{7} and \Re^{6} are each hydrogen atom or optionally substituted $C_{1,g}$ alkyl(as defined above) ring Cy is

EP 1 162 198 A1

(1) $C_{2,0}$ cyclosikyl optionally substituted by 1 to 5 substituent(s) selected from the following group C, group C; hydroxyl group, halogen storn, $C_{1,0}$ sityl and $C_{1,0}$ sitxoxy, (2) $C_{2,0}$ cyclosikonyl optionally substituted by 1 to 5 substituent(s) selected from the above group C, or (3)

- wherein u and v are each independently an integer of 1 to 3,

- (1) C_{B-14} anyl.
 (2) C_{B-6} cycloalkyl.
 (3) C_{B-6} cycloalkenyl or
 (4) haterocyclic group having 1 to 4 hateroatom(e) selected from an oxygen atom, a nitrogen atom and a sulfur
- As and Re are each independently

- (1) hydrogan atom,
 (2) halogan atom,
 (3) optionally substituted G₁₋₆ align (as defined above) or
 (4) -ORse
- wherein Rab is hydrogen atom, C1-8 alkyl or C5-14 aryl C1-8 alkyl, and
- ¥

- (1) hydrogen atom,
 (2) halogon atom,
 (3) cyano,
 (3) cyano,
 (3) cyano,
 (3) cyano,
 (5) amino, C._{1,4} alkanoylamino,
 (6) C._{1,4} alkanoylamino,
 (7) optionally substituted C._{1,4} alkylitas dofined above),
 (9) C._{2,4} alkanyl optionally substituted by 1 to 3 substituent(s) selected from the above group A,
 (9) C._{2,4} alkanyl optionally substituted by 1 to 3 substituent(s) selected from the above group A,
 (9) C.O.G.

- wherein R^{a6} is hydrogen atom or C₁₋₈ alkyt, (10) -CONH-(CH₂)-R^{a10} (10) -CONH-(CH₂)-R^{a10} (10) -CONH-(CH₂)-R^{a10} (10) existituted C₁₋₈ alkyt (as defined above), C₁₋₈ alkoxycarbomy or C₁₋₈ alkanoylamino and 1 is 0 or an integer of 1 to 6, (11) -OR***
- wherein $\mathsf{R}^{\mathsf{alt}}$ is hydrogen atom or optionally substituted $\mathsf{C}_{\mathsf{l-g}}$ altyl (as defined above)
- (12)

wherein wherein and the properties of the proper
--

8	8	â	8	t	8	ig.	<i>8</i> 9	ā	õ	ta.	
(10) (A-(Ch ₂),-CO., (11) (A-(Ch ₂),-O., (12) (A-(Ch ₂),-A-(Ch ₂),- (13) (Ch ₂),-Alper2 (Ch ₂),- whorein Ress is	(6)	(1) a single bond, (2) C _{1-d} elkylene, (3) C _{3-d} elkonylene, (3) C _{3-d} elkonylene, (4) -(CH-J ₀ -P-(CH-J ₀)-, (4) -(CH-J ₀ -P-(CH-J ₀)-, (hersinsfier m and n are each independently 0 or an integer of 1 to 6),	w is an integer of 1 to 3, and Y is	and (p)-(CH ₂)-SO ₂ /NHRe ²⁶ (p)-(CH ₂)-SO ₂ /NHRe ²⁶ (p)-(CH ₂)-SO ₂ /NHRe ²⁶ in hydrogen storn, optionally substituted C ₁₋₄ ally/ (as defined above), C ₆₋₁₄ ary/ optionally substituted by 1 to 5 substituent(a) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substituent(a) selected from the above group B.	(n) -(Crb _a): NHSO ₂ -pa ²⁸ whorein Fa ²³ is hydrogen atom, optionally substituted C _{1,0} alkyl (as dofined above), C _{3,1,4} aryl optionally substituted by 1 to 5 substitutent(s) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substitutent(s) selected from the above group B. (o) -(Crb ₂):-S(O _{1,2} -pa ²³ is as defined above, and q is 0, 1 or 2.	(m) -{CH ₂ } _A NR ^{±28} CO-R ^{±4} whorein R ^{±28} is hydrogen atom, C _{1,4} alkyl or C _{1,4} alkanoyi, R ^{±4} is optionally substituted C _{1,4} alkyl (as defined above), C ₈₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substitutent(s) selected from the above group B.	(1') hydrogen atom, (2') optionally substituted C ₁₋₈ alkyl (as defined above), (2') Optionally substituted by 1 to 5 substituent(e) selected from the above group B, (3') C ₉₋₁₄ anyl C ₁₋₈ alkyl optionally substituted by 1 to 5 substituent(e) selected from the above group B or (5') hotercoycle C ₁₋₈ alkyl optionally substituted by 1 to 5 substituent(e) selected from the above group B,	(k) -(CH ₂) _k -O-(CH ₂) _b -COR-at wherein Rat II of C., allykamine or heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B, and p is 0 or an integer of 1 to 8, (1) -(Ch ₂) _b -N ₁ Agazapaza wherein Razz and Razz are each independently	(b) Trainto-Cycle L., a sity optionally substituted by 1 to 5 substitutently, selected from the above group B, (f) C ₂₋₄ cycloally/ optionally substituted by 1 to 5 substitutently addected from the above group B, or (10°) C ₃₋₄ cycloally/ C ₁₋₄ ally/ optionally substituted by 1 to 5 substitutent(s) selected from the above group B,	(27 optionally substituted C _{1-a} alkyl (as defined above). (37 optionally substituted C _{2-a} alkenyl (as defined above). (47 C _{2-a} alkynyl optionally substituted by 1 to 3 substituent(s) selected from the above group A. (57 C ₆₋₁ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (67 C ₆₋₁ aryl C _{1-a} alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (77 hotorocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B.	EP 1 162 198 A1

(17) hydrogen atom.
(27) optionally substituted C_{1-a} alkyl (as defined above).
(37) C₀₋₁₄ aryl C_{1-a} alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group 8, (47) C₀₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B.

wherein R^{bo} is hydrogen atom, optionally substituted $C_{i,q}$ alkyl (as defined above), $C_{g_{-i,q}}$ anyl optionally substituted by 1 to 5 substitutentle) selected from the above group B or $C_{g_{-i,q}}$ anyl C_{i-q} alkyl optionally substituted by 1 to 5 substitute(s) selected from the above group B, (6) $-COOR^{bo}$ (R^{bo} is as defined above) or

(14) -NRP12CO- (R112 is as defined above), (IS) -CONRP13-(CH₂), (IS) -CONRP13-(CH₂), (IS) -CONRP13-(CH₂), wherein R1213 introduced interm, optionally substituted C₁₋₆ alityl (as defined above) or C₆₊₆ aryl C₁₋₆ alityl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (IS) -CONH-CHR16-(CH₂), (IS) -CONH-CHR16-(IS) as the distribution of the above group B, (IS) -CONH-CHR16-(IS), (IS) -CONH-CHR

(1") hydrogen atom, (2") carboxyl, (3") C₁₋₆ alkyl, (4") -OAbe

wherein R⁵⁶ is C₁₄ alkyl or C₆₊₆ anyl C₁₄ alkyl, or (57 -NHR⁵⁷)
wherein R⁵⁷ is hydrogen atom, C_{1,6} alkyl, C_{1,6} alkanoyl or C₆₊₆ anyl C_{1,6} alkyloxycarbonyl, or R⁶¹⁵ is

wherein n', dng 8; Z and w' are the same as the above-mentioned n, dng 8, Z and w, respectively, and may be the same as or different from the respective counterparts,

(18)-(CH₂) $_{\rm A}$ -NR+12-CHR+13. (R+12 and R+15 are each as defined above), (19) -NR+17SO $_{\rm 2^+}$

8

Ľ,

g

wherein R^{a17} is hydrogen atom or C₁₋₄ alkyl or (20) -S(O)₆-(CH₂)_m-CR^{a13}Ra¹⁶-(CH₂)_n - (e is 0, 1 or 2, IP^{a15} and Ra¹⁶ are each as defined above).

(2) The therapeutic agent of (1) above, wherein 1 to 4 of the G¹, G², G², G², G⁴, G⁶, G⁶, G⁶, G⁶ and G⁶ is (are) a filtragen atom.
(3) The therapeutic agent of (2) above, wherein G² is G(-R²) and G⁶ is a carbon atom.
(4) The therapeutic agent of (2) or (3) above, wherein G² is a nitrogen atom.
(5) The therapeutic agent of (1) above, wherein, in formula [i], the molety

å

8

la a fused ring selected from

EP 1 162 196 A1

(6) The therapeutic agent of (5) above, wherein, in formula [i], the moiety

is a fused ring selected from

(7) The therepeutic agent of (8) above, which comprises a fused ring compound of the following formula [1-1]

wherein each symbol is as dofined in (1),
or a pharmeceutically acceptable salt thereof as an active ingradient.
(8) The therapeutic egent of (6) above, which comprises a fused ring compound of the following formula (1-2)

[-2]

wherein each symbol is as defined in (1), or a pharmaceutically acceptable set thereof as an active ingrodient.

(9) The therapeutic egent of (8) above, which comprises a fused ring compound of the following formula [I-3]

wherein each symbol is as defined in (1),
or a phermaceutically ecceptable eat thereof as an active ingredient.
(10) The therapeutic agont of (6) above, which comprises a fused ring compound of the following formula [1-4]

=

wherein the moiety

wherein R1, R2, R3 and R4 are each independently.

EP 1 162 186 A1

wherein each symbol is as defined in (1), or a pharmaceutically acceptable sat thereof as an active ingredient. or a pharmaceutically acceptable sat thereof as an active ingredient least one of R¹, R², R³ and R⁴ is carboxyl, (11) The therepeutic agent of any of (1) to (10) above, wherein at least one of R¹, R², R³ and R⁴ are as defined in (1), -COORe¹, -CONRe²Ra' or -SO_ARa² wherein Ra¹, Ra², Ra³ and Ra² are as defined in (1). (12) The thorapeutic agent of any of (1) to (11) above, wherein the ring Cy is cyclopentyl, cyclohenyl, cyclohenyl

or tetrahydrothlopyranyl. (13) The therapeutic agent of any of (1) to (12) above, wherein the ring A is
$$G_{b+c}$$
 aryl. (14) A tused ring compound of the following formula (ii)

(1) C _{D-14} aryl,	(3) optionally substituted C ₁₋₆ alkyl (as defined above) or (4) hydroxyl group freg B is	ring A' is a group selected from a group consisting of phenyl, pyridyl, pyrazinyl, pyrimidinyl, pyridazinyl, cy- cibhasyl, cyclohazanyl, furyl and thienyl, A' and Re' are each independently (1) hydrogen storn, (2) helogen storn,	wherein u and v are each independently an integer of 1 to 3.		ring $\mathbf{C}_{\mathbf{y}}$ is $(1) \mathbf{C}_{\mathbf{y},\mathbf{q}} \text{ cyclosikyl optionally substituted by 1 to 5 substituent(s) selected from the following group \mathbf{C}, group \mathbf{C}; hydroxyl group, halogen storn, \mathbf{C}_{\mathbf{t},\mathbf{q}} sitly land \mathbf{C}_{\mathbf{t},\mathbf{q}} sitkoxy, or (2)$	(13). P(=O) (OR=31) ₂ wherein Re31 is hydrogen atom, optionally substituted C ₁₋₂ alkyl (as defined above) or C ₄₋₁₄ anyl C ₁₋₂ alkyl optionally substituted by 1 to 5 audistitutent(s) selected from the above group B, and R ² is hydrogen atom or optionally substituted C ₁₋₂ alkyl (as defined above).	(2) SO ₂ AP. (2) SO ₂ AP. wherein Re ² is hydroxyl group, amino, C ₁₋₆ alkyl or C ₁₋₆ alkylamino	wherein R** is hydrogen atom or hydroxy/ group, (10) -NHAe3 whorein R*5 is hydrogen atom, C _{1,0} alkanoyl or C _{1,0} alkylsulfonyl, (11) -OR a5 wherein R*5 is hydrogen atom or optionally a thetitrated C _{1,0} alkylsulfonyl,	 c) - Connection each independently hydrogen atom, C_{1,6} alkoxy or optionally substituted C_{1,6} alkyl (as defined above), (9) - C(-NR²⁴)NH₂, 	(/) -COOPE* wherein R*1 is optionally substituted Gr ₁₄ alkyl (as defined above) or G ₆₊₁₄ anyl Gr ₁₄ alkyl optionally substituted by 1 to 5 substituent(s) selected from the following group 8, by 1 to 5 substituent(s) selected from the following group 8, group 8, helogen storn, cysno, nitro, Gr ₁₄ alkyl, helogenstad Gr ₁₄ alkyl, Gr ₁₄ alkanoyl, group 8, telegens storn, cysno, nitro, Gr ₁₄ alkyl, helogenstad Gr ₁₄ alkyl, Gr ₁₄ alkanoyl, (Gr ₁₅), -NHSO ₂ RP¹, -(Gr ₂₅), -GOAPP¹, -(Gr ₂₅), -GO	(β) C _{t.,d} alkyl optionally substituted by 1 to 3 substituent(s) selected from the following group A, (g) C _{t.,d} alkylogen atom, hydroxyl group, cerboxyl, amino, C _{t.,d} alkoxy, C _{t.,d} alkoxycerboxyl and C _{t.,d} alkylamino,	(2) Cr. ₄ alkanoyl, (3) carboxyl, (4) cyeno, (5) cyeno, (6) cyeno,
	8	80	49	46	SE S	S		80	ä	ë Bengana	s each	20 CC CC
(1') hydrogen stom,	() -(CH ₂), C(=NR ⁴²⁹)NH ₂ wherein R ⁴²⁵ is hydrogen atom or C ₁₋₂ ellyt. () -(CH ₂), O ₁ Res wherein R ⁴²⁰ is	stituted by 1 to 5 statisticum(s) satisfact from the above group is, as defined above, from (s) of the statistical from the above group is, as defined above, from it is a statistical from the above group is or protection of the statistical by 1 to 5 substitutent(s) selected from the above group is or protection of the statistical by 1 to 5 substitutent(s) selected from the above group 8.	(6") heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B. (6") heterocycle C ₁₋₀ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B.	 (1') hydrogon atom, (2'') optionally substituted C₁₋₈ alkyl (as defined above), (3') C₆₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (4') C₆₋₁₄ aryl C₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above 	(p) -(CH ₂)-COOP=19 wherein Ra19 is hydrogen stom, optionally substituted C ₁₋₈ sliyl (as defined above) or C ₀₋₁₄ snyl C ₁₋₈ alkyl optionally authstituted by 1 to 5 substituent(s) selected from the above group B, (h) -(CH ₂)-CONR437R429 wherein R427 and R438 are each independently,	(2) Ce ₁ , a pri optionally substituted by 1 to 5 substituent(s) selected from the above group B or (3) heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B wherein the nationoxicle group has 1 to 4 hoterostom(s) selected from an exygen storn, a nitrogen storn and a sulfur atom.	wnorain ++** is (1') optionally substituted C ₁₋₆ alkyl (as dafined above),	(c) nice. (d) nice. (e) principle substituted C ₁₋₀ alkyl (as defined above), (f) -(CH- ₂)-COR ¹⁶ . (h) -(CH- ₂)-COR ¹⁶ . (here) hafter each I meane independently 0 or an integer of 1 to 6).	(a) hydrogen stom, (b) habogen stom, (c) ryann	(1) a group selected from the following group D. (2) C ₉₊₄ any optionally substituted by 1 to 5 aubstituent(s) selected from the following group D. (3) C ₉₊₄ expinality optionally substituted by 1 to 5 substituent(s) selected from the following group D. (4) C ₉₊₄ expinality optionally substituted by 1 to 5 substituent(s) selected from the following group D or (5) halarcoyclic group optionally substituted by 1 to 5 substituent(s) selected from the following group D whorein the hotelowing group D whorein the hotelowing group D whorein the hotelowing group D as 1 to 4 heterosiom(s) selected from an oxygen stom, a ntrogen stom and a sulfur stom, group D:	each Z is independently	 C₂₋₈ cyclosityl or heterostom(s) selected from an oxygen atom, a nitrogen atom and a suffur atom,

EP 1 162 186 A1

	8	80	۵		6	ts	સ	25	£0	ā	ē	ts.
(†') hydrogen atom,	(i) -(CH ₂)-C(=NR ⁴³³)NH ₃ wherein R ⁴³³ is hydrogen atom or C ₁₋₆ elkyl, (j) -(CH ₂)-OR ⁴²⁰ wherein R ⁴²⁰ ia	(8") $C_{3,0}$ cyclosity? $C_{1,0}$ sity! optionally substituted by 1 to 5 substituent(s) selected from the above group B.	group B, wherein the heterocycle $C_{1-\delta}$ alityl is $C_{1+\delta}$ alityl substituted by heterocyclic group optionally substituted by 1 to 5 substituent(s) adected from the above group B, as defined above, (7°) $C_{3-\delta}$ cyclosityl optionally substituted by 1 to 5 substituent(s) selected from the above group B, or	(5") heterocyclic group optionally aubstituted by 1 to 5 substituent(s) selected from the above group B. (6") heterocycle C ₁₋₀ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above	(1) hydrogon atom, (2") optionally substituted C _{1.4} alkyl (as defined above), (2") C ₂₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (4") C ₂₋₁₄ anyl C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above croup B.	(g) -(CH ₂)-COORs ⁴⁸ wherein R ⁴⁸ is hydrogen atom, optionally substituted C ₁₋₈ alkyl (as defined above) or C ₈₋₁₈ aryl C ₁₋₈ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (h) -(CH ₂)-CONRSTRAZ (h) -(CH ₂)-CONRSTRAZ wherein R ⁴²⁷ and R ⁴²⁸ are each independently,	girusy o wherein the heterocyclic group has 1 to 4 heterostom(s) solected from an oxygen etom, a nitrogen atom and a suffur atom,	(1) optionally substituted C ₁₄ allyl (as defined above), (2) C ₈₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B or (2) helicocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above	(d) nitro. (e) optionally substituted C ₁₋₀ alkyl (as defined above). (f) -(CH ₂)-COR*1 ⁶ . (hersinative each means independently 0 or an integer of 1 to 6). whomin R*1 ⁶ is	(a) hydrogen atom, (b) halogen atom, (c) cyano,	 a group selected from the following group D. Cop44 any optionally substituted by 1 to 5 substituent(s) selected from the following group D. Cop45 any including production by 1 to 5 substituent(s) selected from the following group D. Cop46 any including substituted by 1 to 5 substituent(s) selected from the following group D or Cop46 any including substituted by 1 to 5 substituent(s) selected from the following group D or Cop46 and produced group based to 4 heterostom(s) selected from an oxygen atom, a nitrogen atom and a sulfur atom, group D. 	each Z is independently

ವ

(2) optionally substituted $C_{1,q}$ silvyl (as defined above). (3) optionally substituted $C_{2,q}$ alixwyl (as defined above). (4) $C_{2,q}$ alixynyl optionally substituted by 1 to 3 substitutint) subscited from the above group B. (5) $C_{2,q}$, anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (9) $C_{2,q}$ anyl $C_{1,q}$ alixyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (2) heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B. (2) heterocycle $C_{1,q}$ alixyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (3) $C_{2,q}$ cyclealixyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (10) $C_{2,q}$ cyclealityl optionally substituted by 1 to 5 substituent(s) selected from the above group B.	riom the above group A, m the above group B, selected from the above selected from the above selected from the above do from the above group bed from the above group ent(e) selected from the
(k) - (CH ₂),-O:(CH ₂),-COR421 wherein R421 is C ₁₋₆ sitylamino or heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B, and p is 0 or an integer of 1 to 6. (f) -(CH ₂),-NR422p422 wherein R422 and R423 are each independently	d by 1 to 6 substituent(e)
(1) hydrogen atom. (2) optionally substituted C ₁₋₄ alityl (as defined above), (3) C ₉₋₁₄ anyl optionally substituted by 1 to 5 substitutent(s) selected from the above group B. (4) C ₉₋₁₄ anyl C ₁₋₆ alityl optionally substituted by 1 to 6 substitutent(s) selected from the above group B or (5) heterocycle C ₁₋₆ alityl optionally substituted by 1 to 6 substitutent(s) selected from the above group B.	m the above group B, selected from the above selected from the above
(m) -(CH ₂) ₁ -NR- ²² CO-R ²⁴ wherein R ²⁵ is hydrogen atom, C _{1-d} alkyl or C _{1-d} atkencyl, R ²⁵ is optionally substituted C _{1-d} alkyl as defined above), C ₂₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B, (n)-(CH ₂) ₁ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₁ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₂ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₂ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₃ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₄ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₄ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₄ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₅ -NHSO ₂ (n)-(CH ₂) ₅ -NHSO ₂ -R ²⁵ (n)-(CH ₂) ₅ -NHSO ₂ (n)-(CH	solided from the above selected from the above selected from the above databove), C _{B+1} and op-18 or helarocyclic group
(0) (Cr ₂)-(c)-(₁ -n-1), (0)-(n-1), (0) (1-n-1), (0) (1	ebove). C ₅₋₁₄ any option- eterocyclic group option-
is an integer of 1 to 3, and is	
(1) a single bond, (2) C _{1-d} alkylene, (3) C _{2-d} alkernylene, (4) -(CH-y ₀ , -)-(CH-y ₀), (morbinatior m and nine each independently 0 or an integer of 1 to 6), (5) -CO>, (6) -CO>, (7) -CONH-(CH-y ₀), NH+, (7) -CONH-(CH-y ₀), NH+,	

EP 1 162 196 A1

```
(8) -NHCONH-.
(10) -C(CH<sub>2</sub>)<sub>1</sub>-CO-.
(11) -C(CH<sub>2</sub>)<sub>2</sub>-CO-.
(11) -C(CH<sub>2</sub>)<sub>2</sub>-CO-.
(12) -C(CH<sub>2</sub>)<sub>3</sub>-CO-.
(13) -C(CH<sub>2</sub>)<sub>3</sub>-CO-.
(14) -C(CH<sub>2</sub>)<sub>3</sub>-CO-.
(15) -C(CH<sub>2</sub>)<sub>3</sub>-(14) by deposition of C<sub>3</sub>-a sity (as defined above).
(2) *Co<sub>2</sub>-a sity (C<sub>1,4</sub> alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B.
(3) *C<sub>2</sub>-1,4 sity (C<sub>1,4</sub> alkyl optionally substituted by 1 to 5 substituted by
```

ú

5

wherein each symbol is as defined in (14), or a pharmacourtically acceptable selt thereof.

(16) The fused ring compound of (14) above, which is represented by the following formula [II-2]

2

ß

wherein each symbol is as defined in (14), or a pharmaceutically accoptable sat thereof.

(17) The fused ring compound of (14) above, which is represented by the following formula (II-3)

8

whoratin each symbol is as doffined in (14).
or a pharmaceuticable acooptable as it hereof.
(19) The fused ring compound of (14) above, which is represented by the following formula [II-4]

EP 1 162 186 A1

wherein each symbol is as defined in (14),
or a pharmaceutically acceptable satt thereot.
(19) The fued ring compound of eary of (14) to (18) above, wherein at least one of A¹, A², A² and A² is carboxyl,
-COOR*¹ or -SO_A*A² wherein A* and A* are as defined in (14), or a pharmaceutically acceptable satt thereot.
(20) The fued ring compound of (19) above, wherein at least one of A¹, A², A² and A² is carboxyl or -COOR*¹

wherein R⁴¹ is as defined in (14), or a phermaceutically acceptable salt thereof.

(21) The fused ring compound of (20) above, wherein R² is carboxyl and R¹, R³ and R⁴ are hydrogen atoms, or a phermacoutically acceptable salt thereof.

(22) The fused ring compound of any of (14) to (21) above, wherein the ring Cy is cyclopernyl, cyclohexyl, cy-cloheyyl or tetrehydrothiopyraryl, or a pharmacoutically ecceptable saft thereof.

(23) The fused ring compound of (22) above, wherein the ring Cy is cyclopernyl, cyclohexyl or cycloheptyl, or a pharmacoutically acceptable saft thereof.

(24) The fused ring compound of any of (14) to (23) above, wherein the ring A' is phanyl, pyridyt, pyrazinyl, pyri-midnyl or pyridazinyl, or a pharmaceurically acceptable salt thereof. (25) The fused ring compound of (24) above, wherein the ring A' is phenyl or pyridyl, or a pharmaceurically ac-

(26) The fused ring compound of (25) above, wherein the ring A' is phenyl, or a pharmaceutically acceptable sattered.

(27) The tused ring compound of any of (14) to (26) above, wherein the Y ts -(CH₂)_m-O-(CH₂)_h. -NHCO₂: -CONH-CHR¹¹, -(CH₂)_h. HR¹²-CHR¹³, -CHR¹³-C

(30) The lused ring compound of any ol (14) to (29) above, wherein the R² is carboxyl, R¹, R³ and R⁴ are hydrogen stores, the ring Cy′ is cyclopentyl, cyclohesyl or cycloheptyl, and the ring A¹ is phenyl, or a pharmacsutbally acselined in (14), or a pharmacoudosily accoptable salt thereof.

ceptable salt thereof.

(31) The fused ring compound of the formula [I] or a pharmaceutically acceptable salt thereof, which is selected from the group consisting of

elhyl 2(4-(3-homophenoxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate (Example 1), 2-(4-(3-homophenoxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylic acid (Example 2),

Ġ

ethyl 1-cyclohexyl-2-(4-hydroxyphony)bonzimidazolo-5-carboxylate (Example 3), ethyl 2-(4-(2-bromo-5-chlorobenzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylate (Example 4), ethyl 2-(4-(2-bromo-5-chlorobenzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylate (Example 4), ethyl 2-(4-(2-(4-chlorophenyl)-6-chlorobenzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylate (Example 4),

2-(4-(2-(4-chlorophenyl)-5-chlorobenzyloxylphenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid (Exemple

othyl 2-(4-(2-bromo-5-methoxybenzyloxy)phenyl]-1-cyclohaxylbenzimidazole-5-carboxylate (Exampte 7), ethyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxylphenyl)-1-cyclohaxylbenzimidazole-5-carboxylate (Ex-

2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)phenyl)-1-cyclohexylbenzimidezole-5-cerboxylic acid (Exam-

othyl 1-cyclohexyl-2-(4-{(E)-2-phenylvinyljphenyl)benzimidezolo-5-carboxylate (Example 10), 1-cyclohexyl-2-(4-{(E)-2-phenylvinyljphenyl)benzimidezole-5-carboxylic acid (Example 11),

8

to 14 hearthmathanil a control to the control of th
2-44-1(4-cenzyroxypheny);caroonylaminojphenyly i-cyclopenyloenzmidazda-o-carodxylid acid (cxamble
2-(4-(14-chlorophenyl)csrbonylaminojphenyl)-1-cyclopenylbenzimidszole-5-csrboxylic scid (Example 59),
1-cyclopentyl-2-[4-(3,5-dichlorophenylcarbonylamino)phonyl]-benzimidazole-5-carboxylic acid (Example 58),
2-(4-(benzenesutionytamino)phenyth 1-cyclopentytbanzimidazole-5-carboxytic acid (Example 57),
2-(4-)3-cnipropentyjoxy)phenyij- i cycupentyjpentymidatoje-progrovyjic acid (Example 68). 2-(4-)apazyjovohanyij-3-ovojennotyjpentyjmidatoja-5-parjonyijo acid (Example 68).
2-(4-(2-chlorobenzyloxy)phonyi]-1-cyclopentylbenzimidezole-5-carboxylic acid (Example 54).
[2-(4-benzyloxyphenyl)-1-cyclopentylbenzimidszol-5-yl]-carbonyleminoscetic acid (Example 53),
61). 1-cyclopentyl-2-(4-hydraxyphenyl)benzímidszole-6-carboxyllo scid (Example 52),
1-cyclopenty-2-(4-((3,6-dimethyl-4-isoxazolyl)methoxylphenyl)-benzimidszote-5-carboxylic acid (Example
) -cyclopenty+2-(4-(4-namaxycenzyloxy)phenylbenzimicazole-o-carboxylic acid hydrochloride (Example 49), 1-cyclopenty+2-(4-(4-pyridyimethoxy)phenyl benzimidazole-6-carboxylic acid hydrochloride (Example 49),
1-cyclopentyl-2-[4- (4-critiuoromethylbenzyloxy)phenylj-benzimidazole-5-carboxyllo acid (Example 47),
2-(4-((2-chlore-5-thlenyl)methoxylphenyl]-1-cyclopontylbenzimidazote-5-carboxylic acid (Example 48),
2-(4-(4-chlorobenzyloxy)phenyli-1-tyclopentylbenzimidezole-5-carboxylic add (Example 45).
2-(4-(4-tort-butylbenzyloxy)phenyi)-1-cyclopentylbenzimidazolo-5-carboxylio acid (Example 43),
6-aultamoyl-2-(4-benzyloxyphanyl)-1-cyclopentylbenzimidazole (Example 42),
o-accryramino-2-(4-benzyloxypneny)-1-cycropentyroenzimioazore (Example 40). 2-(4-benzyloxyphenyl)-1-cycropentyf-5-methanesulfonyl-aminobenzimidazolo (Example 41).
5-amino-2-(4-benzyloxyphenyl)-1-cyclopantybenzimidszole hydrochlonde (Example 39),
2-(4-benzyloxyphonyl)-1-cyclopentyl-5-nitrobenzimidazole (Example 38),
ndo (Example 97).
5-acety-2-(4-benzyloxyphenyl)-1-cyclopentybenzimicazolo (Example 35). 2-/4-benzyloxymhenyl)-1-cyclopentyl-N-/2-dimethylaminosthyl)-benzimidazole-5-carboxamida dihydrochlo-
2-(4-benzyloxyphenyl)-1-cyclopenty-5-(1-hydroxy-1-methylethyl)benzimidazole (Example 35).
2-(4-benzyloxyphenyi)-1-cyclopentyi-N-methoxy-N-methylbenzimidazole-5-carboxamide (Example 34),
erryr z-(e-penzyloxyphenyty- i-cyclopentytentytoenzimidazole-d-carboxamide (Example 33), 2-(4-benzyloxyphenyt)- i-cyclopentyt-N.N-dimethylbenzimidazole-6-carboxamide (Example 33),
2-(4-benzylexyphenyl)-1-cyclopenythenzimidazole (Example 31),
1-cyclohexyl-2-(4-(3-(4-pyridylmethoxy)phenyloxy)phenyl)-benzimidazolo-5-carboxylic acid (Example 30),
ethyl 1-cyclohexyl-2-(4-13-(4-cyridylmethoxy)phenyloxylphenyl)-benzimidazolo-5-carboxylate (Example 29),
othyl 2-(4-(3-acotoxyphenyloxy)phenyl)-1-cyclohoxybenzimidazolo-5-carboxylate (Example 27). ethyl 1-cyclohoxyl-244-(3-hydroxymhenylohoxylnhenylbenzimidazolo-5-carboxylate (Example 28).
2-(4-(3-(3-chlorophanyl)phenoxy)phenyl)-1-cyclohexylbenzlmklazole-5-carboxylib acid (Example 28),
ethyl 2-(4-[3-(3-chlorophenyl)phenoxy)phenyl]-1 -cyclohexylbenzimidazole-5-carboxylate (Example 25).
ethyl 2-(4-benzoylaminopheny)-1-cyclopentylbenzimidazole-5-carboxylate (Example 23),
ethyl 2-(4-eminophemyl)-1-cycloponylbenzimidszole-5-carboxylate (Example 22).
20), athyl 1-cyclopentyt-2- (4-nitrophenylibenzimidazcie-5-carboxylate (Example 21).
2-(4-(bla(3-fluorophonyl)mathoxy); 2-fluorophenyl]-1-cyclohexylbenzimidazola-5-carboxylic acid (Example
etry: 2-14-fats(3-titoropneny)/metnoxy)-2-titoropneny)/-1-cyclonexyloettzimiazole-3-catioxylate (cxembre 19),
ri-2-(2-fluoro-4-hydroxyphenyl)benzimidazole-5-carboxylate (Examplo 18),
i -cyclonexyr.z-q-a-(-a(-a-nuorophenyr)-z-metryr-o-mezolyr-metroxyl[pheny]poetamioazote-o-carovyno ecid (d (Example 17),
late (Example 16),
ethyl 1-cyclohexyl-2-(4-(4-(4-thorophenyl)-2-methyl-5-thiazolyl)methoxylphenyl]benzimidazole-5-carboxy-
2-(4-benzyloxyphenyl)-1-cyclopentybenzimidazole (Example 14), 2-(4-benzyloxyphenyl)-1-cyclopentybenzimidazole-6-carboxamide oxime (Example 15),
2-(4-benzyloxyphenyl)-1-cyclopentybenzimidazole-5-carboxamide (Example 13).
2-(4-benzyloxypheny))-1-cyclopenty/benz/midazele-6-carboxylic acid (Example 12),

EP 1 162 196 A1

2(4-characyclaphaeny)-6-cataonymben/indiacol-1-y)4-d-mithoxyclabnease (Example 63), 2(4-characyclaphaeny)-6-cataonymby-1-cyclaphaeny)-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymby-1-cyclaphaeny-6-cataonymben-2

trans-4-[2-(4-benzytoxyphanyt)-5-carboxybenzimidszol-1-yt]cyclohexan-1-ot (Example 62),

EP 1 162 198 A1

2-(4-(((25)-1-banzenesulfonyi-2-pyrrolidinyi)methoxy[bhenyi]-1-cyclohexy@enzimidazcie-5-carboxyile ecid (Example 183),	
per er), 1-cyclehexy+2-(4-12-(pipendinocarbony/methoxy)phenoxy]-phenyilbenzimidazole-5-carboxyilc acid (Example 192),	8
2-(4-(3-carbamoyi-8-(4-chlorophenyf)bonzyloxy[phenyf)-1-cyclohexy[benzimidazole-6-carboxylicacid (Example 180), 1-cyclohexyl-2-(4-(2-(dimethylcarbamoylmathoxy)phenoxy]-phenyf)benzimidazole-6-carboxylicacid (Example 160),	
יטי, 2-(4-{3-carboxy-6-(4-chlorophony)bonzy/oxy)phonyl}-1-cyctohexy/benzimidazole-6-carboxy/de acid (Example 189)	e e
(Example 169). 1-cycloray/2-(4- [3- (2-pyridylmethoxy)phenoxy]phenyf)-bonzimidazole-5-carboxyfic acid (Example 187). 1-24-(2-(4-chlorophenyf)-5-fluorobenzyfoxy]phenyf)-1-cyclohaxyfbenzimidazole-5-carboxyfic acid (Example 18).	;
io-i; 2-(4-((35)-i-benzyloxycarbonyi-2-pyrrolidinyi)mathoxyjphanyi)-1-cyclohaxylbenzimidazole-5-carboxytic ac-id (Example 185), 2-(2-chioro-4-12-(4-rifiluoromathythenyi)benzyloxyjphanyi)-1-cyclohaxyttenzimidazole-5-carboxytic acid	:
2-(4-(3-acety/amino-6-(4-chlorophany))benzyloxy phany)-1-cyclohaxylbenzimidazole-5-carboxylic acid (Exemple 183), 2-(4-(2-(4-carboxyphanyi)-5-chlorobanzyloxy phanyi)-1-cyclohaxylbenzimidazole-5-carboxylic acid (Example 1974)	8
- I-oydolnosyl-2-(4-13-(2-propyryloxy)phenoxy)phenyl)enzimidazole-3-carboxylic acid (Example 177), 1-oydolnosyl-2-(4-13-(3-pridy)methoxy)phonoxy)phonyl-poralimidazole-3-carboxylic acid (Example 178), 2-(4-tenzyloxy/3-methoxypheny)-1 - (ydolnay)benzimidazole-3-carboxylic acid (Example 178), 2-(4-(2-tenzyloxy)-3-methoxybenzyloxy)phenyl-1 - (ydolnay)benzimidazole-3-carboxylic acid (Example 180), 2-(4-(2-tenzyloxy)phenyl-1-tenzyloxy)phenyl-1 - (ydolnay)benzimidazole-3-carboxylic acid (Example 181), 2-(4-2-(4-chlorophenyl)-3-nilrobenzyloxy)phenyl-1 - (ydolnay)benzimidazole-3-carboxylic acid (Example 182),	8
2-44-2(scan)-4-piparidyi)methoxyjphenoxyjphenyi)-1-cyclohaxyibenzimidazcie-6-carboxyiic ecid (Exam- ple 175). 2-44-2-(1-eceny-4-piparidyi)methoxyjphenoxyjphenyi)-1-cyclohaxyibenzimidazcio-6-carboxyiic ecid (Exam- ple 176).	8
171), 2-(4-(4-bonzyloxyphonoxy)-3-fluorophonyl)-1-cyclohoxybenzimidazolo-5-carboxylic acid (Example 172), 2-(4-(2-bronno-5-chlorobenzyloxyphonyl)-1-cyclohoxybenzimidazole-5-carboxylic acid (Example 173), 2-(4-(2-bronno-5-chlorobenzyloxyl)-2-fluorophonyl)-1-cyclohoxybenzimidazole-5-carboxylic (Example 174)	25
(Chairpier 107). 2-(4-13-chloro-4): Chilenyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Exemple 168). 2-(4-13-chloro-6-(3-chlorophenyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 168). 2-(4-3-chloro-6-(3-pyridyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 2-(4-3-chloro-6-(3-pyridyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 2-(4-3-chloro-6-4-4-fluorophenyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 2-(4-3-chloro-6-4-fluorophenyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 2-(4-3-chloro-6-4-fluorophenyi)benzyloxyjphonyi)-1-cyclohoxyfbonzimidazole-5-carboxylic acid (Example 170).	8
2-(4-1((2S)-1-bonzyl-2-pyrrolidinyl)methoxylphenyl)-1-cyclohoxyl-benzimidazolo-5-carboxylic acid hydrochlo- ride (Example 165), 2-(4-13-chloro-6-(4-methythiophenyl)benzyloxylphenyl)-1-cyclohaxylbenzimidazole-5-carboxylic acid (Exam- ple 166), 2-(4-3-chloro-6-(4-methanesulfonylphenyl)benzyloxylphenyl)-1-cyclohaxylbenzimidazolo-5-carboxylic acid	ä
162). 1-cyclbhaxyl-2-(4-(3-(4-methyl-3-pentenyloxy)phenoxy)phenyl)-benzimklazole-5-carboxylic edid (Example 163). 1-cyclohoxyl-2-(4-(3-(3-methyl-3-butenyloxy)phenoxy)phenyl)-benzimklazole-5-carboxylic edid (Example 1-cyclohoxyl-2-(4-(3-(3-methyl-3-butenyloxy)phenoxy)phenyl)-benzimklazole-5-carboxylic edid (Example 194).	ő
1-cyclohoxyk-244-[2-(4-pbortdyimethoxy)phenoxy]phenyl)-benzimidezote-6-carboxytic edd hydrochloride (Example 160), 1-cyclohoxyk-24-(4)-(4-pbortdyimethoxy)phenoxy]phenyl)-benzimidezote-5-carboxytic edd hydrochloride (Example 161), 2-(4-((2R)-2-acetylamino-2-phenylethoxy)phenyl)-1-cyclohoxylbenzimidezote-5-carboxytic edd (Example 2-(4-((2R)-2-acetylamino-2-phenylethoxy)phenyl)-1-cyclohoxylbenzimidezote-5-carboxytic edd (Example 2-(4-((2R)-2-acetylamino-2-phenylethoxy)phenyl)-1-cyclohoxylbenzimidezote-5-carboxytic	4

	(Example 229),
	2-(4-(2- (4-chbrophenyi) -6-ethoxycarbonybenzyloxy phenyi)-1-cyclohexylbenzimidezole-6-cerboxylic ecid
	2-(4-((2S)-1-(4-dimethylcarbemoylphenyl) -2-pyrrolidinyl)-methoxylphenyl)-1-cyclohexylbenzinidezole-
	2-(4-(4-chlorobonzyloxy)piperidinojphonyli-1-cyclohexylbenzimidazole-5-carboxylic add (Exemple 22s), 2-(4-(3)(2-chloro-4-pyridy)jmethoxyjphenoxyjphenyl)-1-cyclohexylbenzimidazole-5-carboxylic add (Exemple 22s), ne 2777
	2-(4-{4-carbamoyi-2-(4-chlorophenyi)benzyloxy phenyi)- i-cyclohexylbenzimidazole-5-carbaxyiic acid (Exam- ple 225),
	2-(4-1-(4-chlorobenzyl)-4-piperdyloxy[phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid (Example 223), 2-(4-3-(4-chlorobenzyloxy)piperdino)phenyl}-1-cyclohexylbenzimidazole-5-carboxylic acid (Example 224),
	1-cyclohexyl-2-(4-(3-(3.5-dichlorophenyl)phenoxylphenyl)-benzimidazole-5-carboxylic scid (Exemple 222),
	1-cyclohoxyl-2-(4-(3-((2,4-dimethyf-6-thlazolyl)methoxylphenoxyl-phenyl)bonzimidazolo-6-carboxylio acid
	1-cyclohexyl·2-[4-[3-[(2-methyl-4-thiszoly))methoxy]phonoxy]-phony]bonzimidszolo-5-carboxylia ecid (Ex- ample 220),
	(Example 2.0). 2-(4-{1-(4-chlorobenzy)}-3-piperidyloxy)phenyi)-1-cyclohexybenzimidazole-5-carboxylio acid (Example 219),
	2-(4-((4-(4-chlorophenyl)-2-methyl-6-thlezofyl)methoxylphenyl)-1-cyclohexylbonzimidezole-5-carboxylic acid
	1-cyclohoxy/-2-(4-(4-(4-methanesulfonytphenyl)-2-mothyr-5-thiazolyty/jmethoxy/phenyl)benzimidazole-
	2-(4-(3- (4-chloropheny)) phenoxy(pheny))-1 -cyclonexy(benzimicazole-6-carboxyilo acid (Example 216), 1-cyclohexy(-2-(4-13-(4-methoxypheny))phenoxy(pheny))-benzimidazole-6-carboxyilo acid (Example 216),
	1-cyclohexyl-2-(4-(3-(3-pyridyi)phenoxyjphenyljbenzimidszole-5-carboxylic acid (Example 214),
	2-[4-[3-(4-tert-buty/benzy/oxy/phenoxy/phenyi]-1-cyclohexy/benzimidazole-5-carboxy/lo ecid (Example 212), 2-[4-[3-(2-chlorobenzy/oxy/phenoxy/phenyi]-1-cyclohexy/benzimidazole-5-carboxy/lo ecid (Example 213),
	1-cyclohexyl-2-(4-(3-(1-methyl-4-pipendy))methoxy)phenoxy)-pheny)joonzimida2oie-5-carooxyiic acid (Ex- ample 211),
	210),
	209), 1-cyclohoxyl-2-44-13-44-trifiuoromethylbenzyloxylohenoxylohenyl-benzimidazote-5-carboxylic acid (Example
	1-cyclohexyl-2-(4-(3-(4-tetrahydropyranyloxy)phenoxy)phenyl)-benzimidazole-5-carboxylic ecid (Example
	1-cyclohexyl-2-(4-(2-(4-chlorophenyl)-3-nltrobenzyloxylphenyl)-benzimidazolo-8-carboxylio acid (Example 208).
	(Example 200), 2-(4-[bis(3-fluoropheny)]methoxyjphenyl]-1-cyclohexylbenzimidazole-5-carboxylic acid (Example 207),
	2-[4-[(5-(4-chlorophenyl)-2-methyl-4-thlazolyl)methoxylphenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid
•	2-[4-{((≥s)-1-(4-acetylaminophonyl)-2-pyrrollamyl)methoxyl-phenyl)-1-cyclonexyloenzimieszole-e-≂arouxyl- ic acid (Example 205),
	chloride (Example 204),
	1-cyclohexyl-2-(4-(((2S)-1- (4-nitrophenyl) -2-pyrrolldinyl)-methoxylphenyljbenzimidazole-6-carboxylic acid
	2-(4-(3-(4-chlorobenzyloxy))phenoxy)phenoxy)phenoxylphenoxylphenzimidazole-5-carboxylic acid (Example 201). 1-cyclohexyl-2-(4-13-(4-fluorobenzylpxy)phenoxylphenyl1-banzimidazole-5-carboxylic acid (Example 202).
	2-(4-(3-chlorobenzyloxy)phenoxylphenyl)-1-cyclohexylbenzimidszole-5-csrboxylic acid (Exemple 200).
	1-cyclohexyl-2-4-([2-methyl-5-(4-chlorophenyl) -4-oxazolylj-methoxylphenyljdenzimtgazole-b-cerdoxylid ac- id (Example 199),
	acid (Example 198),
	pie 197), 1-avdohexvi-2⊣443-{(1-methanesutfonvi-4-abeddvtimethoxy)-phonoxylphenvtibenzimidazolo-5-carboxylic
	1-cyclohexyl-2-(4-(3-(pipendinocarbonylmethoxy)phenoxy)-phenyl)benzimidazole-5-carboxylic acid (Exam-
	1-cychroxy-2-(4-(3-(dimethylcarbemoylmethoxy))phenoxy)-phenyljbenzimidazole-5-carboxylic acid (Exam-
	2-(4-(2-(4-carbamoylphanyl)-5-chlorobenzyloxylphanyl)-1-cyclohexylbanzimidazola-5-carboxylic acid (Exam- nia 1947)
	2-(4-{((2S)-1-banzayl-2-pyrrolidinyi)mathexyiphenyij-1-cyclahoxyibenzimidazale-5-carbaxyila acid (Example 194).

EP 1 162 198 A1

2-(4-(2-(4-chloropheny))-5-methoxybenzyloxy]phenyl]-1-cyclohexylbenzimidazole-4-carboxylic acid (Exam-	
1-cyclohexyi-2-[4-[3-carboxy-5-[4-pyrfdylmethoxy)phenoxy]-phenyi]benzimidazole-5-carboxylic acid dihydro- chloride (Example 260),	
ecid dihydrochlorido (Example 259),	ä
2-(4-(2-(4-chlorophonyl)-6-mothoxybenzyloxy)phenyl)-3-cyclohoxylbenzimidazole-4-carboxylic acid (Example 256),	
(Example 250), 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxylphenyl)-1-cyclohexylbenzimidazote-5-euftenic acid (Example 257)	50
acid hydrochlorida (Example 255), 1-cyclohaxyl-2-42-fluoro-4-(4-fluoro-2-(3-fluorobanzoyl)-benzyloxy phenyl]benzimidazole-5-carboxylic acid	
1-cyclohexyl-2-(4-(4-cerboxyphenyl)-2-methyl-5-thtezolyf)-methoxy/phenyf)benzimidazole-5-cerboxylic	i
1-cyclohexyl·2-(4-((4- (4-fluorophenyl) -2-hydroxymethyl-5-thlazolyfjmethoxylphenyljbenzimidazole-5-car- boxylo ecid (Exemple 264)	Ġ.
2-(2-(2-biphenyy)bxymethy):5-thienyi]-1-cyclohexyteenzimidazole-5-carboxyila acid (Example 252), 2-(2-(2-biphenyy)bxymethy):5-tury):1-cyclohexyteenzimidazole-5-carboxyila acid (Example 253),	
oscewe (cxemple zev), 2-(4-benzyloxycyclohexyl)-1-cyclohexylbenzimidszole-5-carboxylic sold hydrochloride (Example 251),	ŧ
2-[4-[2-(4-chlorophenyl)-5-surlamoylbenzyloxy]phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid trifluor-	\$
2-[4-[3-(tert-but/visulfamoyi)-6-(4-chlorophenyi)benzyloxyi-phenyi)-1-cyclohexyibenzimidazolo-5-carboxyib acid (Framola 249).	
hydrochloride (Example 248),	;
yisto (example 247). 2.14.13.14hilomothenvil-5-methyloethennykhenvilotyvibhenvilot-cyclobaryimidevola-5-rethonvilotecid	t
ייטרויטייטים (באמויקים פייטר, methy (24-4-chloropheny) ל-methy (24-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-	
mothyl 2-(4-(5-carboxy-2-(4-chlorophenyl)benzyloxy)phenyl)-1-cyclohoxy/benz/midazole-5-carboxylato hy-	
boxylate (Example 245),	30
empte 244). methyl 2-(445-tort-butoxycarbonyl-2-(4-chtorophenyl)benzyloxyl-phenyl)-1-cyclohoxylbenzimidazole-5-car-	
methyl 2-14-(2-bromo-6-tent-butoxycarbonylbenzyloxy)phenyl]-1-cyclohexylbenzimidazole-6-carboxylate (Ex-	
ony i arteto-ferondropheny pytanina y menony prienty i royalenony centrimez die okazioany idae.	ŧ
chloride (Example 242),	2
2-(4-(2-(4-chloropheny))-5-methoxybenzyloxy)phenyl)-1-cyclohoxyl-benzimidazole-5-carboxylic ecid hydro-	
methyl 2-(4-(2-(4-chlorophonyl)-5-methoxybenzyloxylphenyl)-1-cyclohexybenzimdazole-5-carboxylate (Ex-	
	8
239). 2-14-14-13-chibrophenyî-8-eyrimidinyloxylphenyî-1-eyriphexyîbenzimidezolo-5-carboxylic edit (Example	
1-cyclohexyl-2-(4-(4-(4-pyridyimethoxy)-8-pyrimidinyloxy)phenyl)-benzimidazolo-6-carboxylic acid (Example	
ample 237).	15
2-(4-(2-(4-chlorophenyl)-4-(5-tetrezolyl)benzyloxylphenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid (Ex-	
2-(4-(2-(3-chloropheny))-4-methylamino-1,3,5-triazin-6-yloxy]phanyl}-1-cyclohexylbenztmidazole-5-carboxy/- io acid triftuoroseolate (Example 238).	
235),	
cnionoe (Example 234). 2-44-(3-(4-chloropheny)-2-pyridyi}mathoxyjphenyi}-1-cyclohexythenzimidazole-5-carboxytic acid (Example	ō
2-(4-(2-(4-chlorophenyl)-3-pyridyl)methoxylphenyl]-1-cyclohexylbenzimidazole-5-carboxylic ecid dihydro-	
z-(((
ld (Example 232).	Ça
2-(4-(2-(4-chlorophenyl)-5-dimethylcarbamoylbenzyloxy phenyl)-1-cyclohexylbenzimktazole-5-carboxylic ac-	
1-cyclohexyl-2-(4-1(4-(4-dimethylcarbamoytpherryl)-2-methyl-5-thiazolyl)methoxylpherryl)benzimidazole-	
1-cyclohexyl-2-(4-(3-trifluoromethylphenoxy)phenyfibenzimidazole-5-carboxytic acid (Example 230),	

ß

EP 1 162 196 A1

methyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxyjphenyl)-1-cyclohoxyl-1H-indole-5-carboxylate (Exam-	
2-{4-{2· (4-chlorophenyl) -5- (N-benzyl-N-methylcarbameyl)-benzylexy}-2-fluorophenyl}-1-cyclohexylbenzim- idezele-5-carboxylic acid hydrochlonde (Example 319),	
zole-5-carboxylic acid dihydrochloride (Example 318),	8
zole-6-carboxylic add hydrochlorda (Example 317), 2-14-12-14-chlorophenyll-5-14-pyrigytmothytesrbamoylbenzyloxyl-2-illusrophenyll-1-cyclohexylbonzimida-	
boxylic acid hydrochloride (Example 316), 2-(4-(2-(4-chlorophenyl)-5-(cyclohexylmothylcarbamoyl)benzyloxyl-2-fluorophenyl-1-cyclohoxylbcnzimida-	
2-[4-[2-(4-chlorophenyl)-5-(benzylcarbamoyl)benzyloxyl-2-fluorophenyl)-1-cyclohexylbenzimidazob-5-car-	50
z-(4-to-chloro-z-(4-byrrayi)zenzyłożyj-z-tiuoropnenyi)- i -cyclonexylbenzunidazole-5-carboxylic ecid hydro- chloride (Example 315),	
5-carboxylete (Example 314),	
mothy/ 2-(4-chlorophenyl)-5-(methylcarbamoyl)benzyloxy)-2-(luorophenyl)-1-cyclohexylbenztmidazole-	
314). 2-14-(phenyl-3-pyridylmathoxy)-2-fluoroophanyi)-1-cyclohaxyibanzimidazolo-5-carboxyiic acid (Example 313).	à
2-(4-fbis(4-carboxyphenyf)methoxy]-2-fbiorophenyf]-1-cyclohexyfbenzimidazole-5-carboxyfic acid (Example	
chloride (Example 311),	
arochiondo (Exempio 310), 2-14-12-14-chtorophenyt-5-methoxybenyytthiotohenyt-1-aydohenythenytmidazoh-5-z-s-havytic edit hydro-	å
2-(4-(5-(4-chlorophenyl)-2-methoxybenzylsulfonyl)phenyl)-1-cyclohexylbenzimidazole-5-carboxylib acid hy-	;
drochloride (Example 309),	
SUB),	
2-(4-(5-emino-2-(4-chlorophenyi)benzyloxy)phenyi)-1-cydohexyibenzimidezele-5-carboxylic acid (Example	£
boxyle acid (Example 307),	
2442-44-cartemoviohenvi)-5-4 imanibylcartemovijbenvijovi-phenvijo teoriohenvijonijenvijonijenvijo-8-cer	
2-(4-)2-(4-cardoxypnenyi)-o-meinoxybenzyloxyjpnenyi)-1-cyclonexylbonzimidazolo-5-carboxylic acid (Exam- olo 308).	
(Example 305).	8
2-(4-(6-carboxy-2-(4-chlorophenyi)benzyloxy)-2-lluerophenyi)-1-cyclohexylbenzimidazole-6-carboxylic acid	
20	
eodium 2-44-(2-44-chlorophenyl)-5-(dimethylcarbamoyl)banzyloxyl-2-fluorophenyl-1-cyclohexylbenzimida-	
Zolo-S-carboxylato (Example 303)	
口のけい クルル・グメルトからてのからせい かんぱ かっかい アップ・アップ・アップ・アップ・アップ・アップ・アップ・アップ・アップ・アップ・	è
sowieri z fyzychienymoniemożyj z neodpienyj i zycionexyjodnizmigazoje o zarożyjajo (Exam- nie pos	
endium Odedoshinovkodshinovkovkovkovkovkovali (example sovie a populari (Evan.	
2-(4-(Dis(4-dimethyloarbarroy/phenyl)methoxy)-2-fluorephenyl)-1-cyclohexylbenztmidazele-5-carboxylic acid	
2-(4-(bis(3-pyridy)/momoxy)-2-(luorophenyl)-1-cyclohaxylbenzimidazole-5-carboxylic acid (Example 300).	á
ride (Exemple 288),	:
2-(4-(2-(4-chiarophenyi)-5-cyanabenzyloxyjphenyi)-1-cyclohexylbenzimidazole-5-carboxylic acid hydrochio-	
ampie 298), ampie 298),	
10 (4)	ě
2-(4-(2-(4-chlorophenyl)-5-methylthlobenzyloxyjphenyl)-1-cyclohexytbenzimidazole-5-carboxylic acid (Exam-	:
296),	
2/4/2-:3-septo-youther-chtoroben zyloxylbhenyl)-1-cycloberylbenzimidezole -5-cephovylic edit (Evennie	
kadrochlonde (Examole 295).	;
grochlorido (Exemplo 284),	8
2-(4-(3-chloro-8-(4-hydroxymethylphenyl)benzyloxy)phenyl}-1-cyclohexylbenzimidazole-5-cerboxylic ecid hy-	
(Example 283).	
2-(4-(2-(-4-(2-carboxyothyt) phenyt)-5-chlorobanzytoxy] phenyt]-1-cyclohexytbenzimidazolo-5-carboxytic acid	
zole-6-carboxylic add hydrochloride (Example 292).	5
o-caroxylic acid hydrochloride (Example 291).	
2-(4-{2-(4-chlorophenyl)-5-thlomorpholinocerbonylbenzylaxy]-2-fluorophenyl)-1-cyclohexylbenzimidezole-	
boxylic acid hydrochlorida (Exampio 290),	

pie 501), 2-(4-(2-(4-chlorophenyi)-5-methoxybenzyloxyjphenyi)-1-cydohexyl-1H-indole-5-carboxyllo acid (Example

2-(4-benzyloxyphenyl)-1-cyclopentyl-IH-indolo-5-carboxylia acid (Example 603), ethyl 2-(4-benzyloxyphenyl)-1-cyclopentheaylimidazol; 2-alpyridine-7-carboxylias (Example 601), ethyl 2-(4-benzyloxyphenyl)-3-cyclohaylimidazol; 2-alpyridine-7-carboxylia etaid (Example 602), end 2-(4-f2-(4-chlorophenyl)-5-melhoxybrenzyloxy)phenyl)-3-cyclohexyl-3H-imidazol(4,5-b)gyridine-6-carboxylic 2-(4-f2-(4-chlorophenyl)-5-melhoxybenzyloxy)phenyl)-3-cyclohexyl-3H-imidazol(4,5-b)gyridine-6-carboxylic

(32) A pharmaceutical composition comprising a fused ring compound of any of (14) to (31) above, or a pharmaceutically acceptable carrior.

(33) A hepatitia C virus polymerase inhibitor comprising a fused ring compound of any of (1) to (31) above, or a

pharmaceutically acceptable salt thereof, and a pharmaceutically acceptable carrier

(34) An anti-hepatitis C virus agent compristing a fused fing compound of any of (1) to (31) above, or a pherma-countiestly acceptable satt thereof, and a pharmaceutically acceptable carrier.

(35) A therapoutic agent for hepatitis C comprising a fused ring compound of any of (14) to (31) above, or a pharmaceutically acceptable satt thereof, and a pharmaceutically acceptable carrier.

(35) A method for treating hepstitis C, which comprises administering an effective amount of a fused ring compound of the above-mentioned formula (i) or a pharmacoutically accopitable satt there).

(37) A method for inhabiting hepstitis C virus polymeraes, which comprises administering an effective amount of a lused ring compound of the above-mentioned formula (i) or a pharmacoutically acceptable satt thereof.

(38) Use of a fused ring compound of the above-mentioned formula (i) or a pharmacoutically acceptable satt thereof.

8

(39) Use of a fused ring compound of the above-mentioned formula [1] or a pharmaceutically acceptable salt thereof for the production of a pharmaceutical agent for treating hepatitis C

(40) A pharmacourtical composition for the treatment of hepatits C, which comprises a fused ring compound of the above-mentioned formula (1) or a pharmacoutically acceptable can thereof, and a pharmacoutically acceptable for the production of a hepatitis C virus polymerase inhibitor.

(41) A pharmaceutical composition for inhibiting hepatitis C virus polymerase, which comprises a fused ring compound of the above-mentioned formula [i] or a pharmaceutically acceptable self thereof, and a pharmaceutically

(42) A commercial package comprising a pharmaceutical composition of (40) above and a written matter associated therewith, the written matter stating that the pharmaceutical composition can or should be used for treating hepatitis

atitis C virus polymerase. (43) A commercial package comprising a pharmaceutical composition of (41) above and a written matter associated therewith, the written matter stating that the pharmaceutical composition can or should be used for inhibiting hep-

ŝ

벊

ક

è

Ġ

ક alky having 1 to 4 carbon atoms. Examples thereof include fluoromethyl, difluoromethyl, trifluoromethyl, bromomethyl, chloromethyl, 1.2 citchtoromethyl, 1.2 citchtoromethyl, 1.2 citchtoromethyl, 2.2 citchtoromethyl, 1.2 citchtoromethyl, 2.2 citchtoromethyl, at group B. (1992). The halpoperated C., alkylide particularly preferably trifluoromethyl at group B. (1992). The C., alkylide is streight chain alkylidene having 1 to 6 carbon atoms, and is exemptified by methylono, orbitylane, trimethylene, tetramethylene, pentamethylene or havarenthylene.

(1994) The C., alkylidene is streight chain alkylidene having 2 to 6 carbon atoms, and is exemptified by vinylene, to 1994). The C., alkylidene is streight chain alkenylone or baylane at Y. (1994). [0038] The halogensted C_{14} alkyl is the above-defined C_{14} alkyl except that it is substituted by the above-defined halogen atom. Preferably, it is halogenated alkyl wherein the alkyl molety thereof is straight chain or branched chain

g

The C_{1-A} alkylane is preferably matriylane or ethylene at Y. The C_{2-A} alkenylane is straight chain alkenylane having 2 to 8 carbon atoms, and is exemplified by vinylane føns, 1-butsnylane, 1,3-butsolanylane and the like.

EP 1 162 196 A1

alkoxy wherein the alkyl molety thereof is straight chain or branched chain alkyl having 1 to 4 carbon atoms. Examples thereof include methoxy, ethoxy, propoxy, isopropyloxy, butoxy, isobutyloxy, ten-butyloxy, pentyloxy, haxyloxy and the [0043] The $C_{2,6}$ alkenylens is proforably vinylens at Y.

[0044] The $C_{1,6}$ alkoxy is alkyloxy wherein the alkyl moisty thereof is the above-defined $C_{1,6}$ alkyl. Proforably, it is

[0045] The C_{1-d} elikory is particularly preferably mathoxy at R^{a2}, R^{a3}, group A and group C.
[0045] The C_{1-d} elikancy is elikycarboxyl wherein the alkyl molety thereoil is the above-defined C_{1-d} elikyl. Preferably,
it is elikancyl wherein the alkyl molety thereoil is straight chain or branched chain elkyl having 1 to 4 carbon atoms.

Examples thereoil include acetyl, propietryl, butyryl, isobutyryl, phatleyl and the like.

õ

ä [0047] The C₁₋₆ alkanoyl is particularly preferably acetyl at R1, R2, R3, R4, R4³, R4³, R4³ and group B. [0048] The C₁₋₆ alkoxycarbonyl is sityloxycarbonyl wherein the alkoxy molety thereol is the above-defined C₁₋₈ alkoxy. Preferably, it is alkoxycarbonyl wherein the sityl molety thereol is straight chain or branched chain alkyl having alkoxy. Preferably, it is alkoxycarbonyl wherein the sityl molety thereol is straight chain or branched chain alkyl having 1 to 4 carbon storns. Examples thereof include methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropyloxycar-bonyl, butoxycarbonyl, labbutyloxycarbonyl, tarl-butyloxycarbonyl, pentyloxycarbonyl and the like. [0049] The C_{1-q} sikkoxycarbonyl is particularly preferably methoxycarbonyl or ethoxycarbonyl at Pari'e and group A. [0050] The C_{1-q} sikylamino is alkylamino or disikylamino wherein the alkyl molety thereof is the above-defined C_{1-q} alkyl. Preferably, it is alkylamino or disikylamino wherein the alkyl molety thereof is straight chain or branched chain

8 alkyl having 1 to 4 carbon atoma. Examples thereof include methylamino, ethylamino, propylamino, isopropylamino, butylamino, isopropylamino, butylamino, isobutylamino, terboutylamino, pentylamino, havylamino, dimethylamino, distrylamino, methylathylamino, N-isopropyl-N-isobutylamino and the like.

[0051] The C₁₋₉ alkylamino is particularly preferably methylamino at R^{ay}, and particularly preferably dimethylamino at R^{act} and group A.

8 pivaloylamino and the like. [0053] The C₁₋₀ alkanoyli [0052] The C₁₋₄ elkancylamino is elkylcarbonylamino whorein the elkancyl molety thereof is the above-defined C₁₋₉ elkenoyl. Preferably, it is elkylcarbonylamino wherein the elkyl molety thereof is streight chain or branched chain alkyl having 1 to 4 carbon etoms. Examples thereof include acetylamino, propionylamino, butyrylamino, isobutyrylamino,

ä (2003) The C₁₋₄ alkanoylamino is particularly preferably acetylamino at X and Fe¹⁰. [2004] The C₁₋₄ alkylutforty is altylautforty whorein the alkyl moioty thereof the above-defined C₁₋₄ alkylutforty is altylautforty whorein the alkyl moioty thereof is artight chain to branched chain alkyl having 1 to 4 carbon orably, it is altylautforty whorein the alkyl moioty thereof is artight chain to branched chain alkyl having 1 to 4 carbon orably, it is altylautforty whorein the alkyl moioty thereof is a stage that alkylutforty, bropylautforty, isopropylautforty, bropylautforty, isopropylautforty, bropylautforty, bro

ysautiony/, terbutysutiony/, pentysutiony/, hexylautiony/ and the title. [0055] The C₁₋₄ alkylautiony/ is pacticularly prolonably methylautiony/ at X and Re³. [0056] The C₀₋₄ any/ is aromatic hydrocarbon having 80 of 4 carbon atoms. Examples thereof include pheny/, naph-thyl, incleny/, azuleny/, fluoreny/, phonanthy/ and the like. [0057] The C₀₋₁₄ any/ is preferably pheny/ or naphthyl, penfecularly preferably pheny/ at the ring A, ring A', ring B and

ring B The C_{6.14} anyl is preferably phenyl or naphthyl, particularly preferably phenyl at the ring A, ring B and

ၾ

[0088] The C₃₆ cycloalkyl is saturated cycloalkyl having 3 to 8, protorably 5 to 7, carbon atoms. Examples thereof include cyclograpyl, cyclobutyl, cyclopontyl, cyclohavyl, cyclohaptyl and cyclooctyl.

[0059] The C₃₋₆ cycloalky(is particularly preferably cyclohexyl at the ring A, ring A', ring B and ring B'. [0050] The C₃₋₆ cycloalkenyl is cycloalkenyl having 3 to 8, preferably 5 to 7, carbon atoms and has at least 1, pref

څ

erably 1 or 2, double bond(s). Examples thereof include cyclopropenyi, cyclobutenyi, cyclopentenyi, cyclopentadienyi, cyclopentenyi, cyclopentadienyi, cyclopentenyi, 2,4-cyclohaxadien-1-yi, 2,5-cyclohaxadien-1-yi, cyclohepienyi and cyclocdenyi and the like, but do not include aryl (e.g., phenyl) or completely saturated cycloalkyl

à [0081] The Cae cycloalsony is preleably cyclobaxony at the ring A and ring A.

[0082] The heterocyclic group has, as an atom constituting the ring, I to a heteroatom(s) selected from an oxygen
storn, a nitrogen atom and a suffur storn, basides a carbon atom, and includes saturated ring and unsaturated ring,
monocyclic ring and fused ring having the number of ring atom constituting the ring of 3 to 14.

[0083] The heterocyclic group as monocyclic ring includes, for example, pyrioty, pyrimidiny, pyridaziny,
1,3.5-rinaziny, pyrroyl, pyrazoly, imidazoly, 1,2.4-riazoly, tetrazoly, thienyl, hury, exazoly, teoxazoly, thiazoly, tetrazoly, twenty, hury, exazoly, teoxazoly, thiazoly, terrazoly.

50 thiazolyi, thiadiazolyi, pyrrolinyi, pyrrolidinyi, imidazolidinyi, piperidyi, piperazinyi, morpholinyi, thiomorpholinyi, tetrahy

[0064] Examples of the heterocyclic group as a fused ring Include quinolyt, Isoquinolyt, quinazolinyt, quinoxalyt, phthatazinyt, cinnolinyt, naphthyridinyt, 5,8,7,8-tetrahydroquinolyt, Indolyt, benzinidazolyt, Indolinyt, benzoturanyt, benzo

thereot include pyridyl, pyrazinyl, pyrimidinyl, pyridazinyl, 1,3,5-thazinyl, pyrrolyl, pyrazolyl, imidazolyl, 1,2,4-triazolyl Istrazolyl, ithlenyl, furyl, oxazolyl, isoxazolyl, ithazolyl, isothlazolyl, thiadiazolyl, pyrrolidinyl, pipendyl, pipenazinyl enc [0065] Preferably, it is a heterocyclic group which is a 5-membered or a 6-membered monocyclic group. Examples

[0889] The heterocyclic group is preferably pyridyf, pyrazinyf, pyrimidinyf or pyridazinyf which is an aromatic group, particulary perforably pyridy at the ring A and ring A.
[0887] The heterocyclic group is particularly preferably pyridyf, pyrazinyf, pyrimidinyf, pyridazinyf, 1,3.5-triazinyf, pyrrodyf, pyrazobyf, imidazobyf, 1,2.4-triazobyf, isterazobyf, ithlanyf, furyf, oxezobyf, isoxazobyf, is azolyl, which is an aromatic group, at the ring B and ring B'. More preferably it is pyridyl or thiazolyl, most preferably

õ propyl, 2-phenytpropyl, 4-phenylbutyl and the like. [0089] The $C_{6,14}$ aryl C_{1-6} alkyl is particularly pro [0088] The $G_{\Phi/4}$ any $G_{1-\theta}$ alky its anylalky wherein the sikyl molely thereof is the above-defined $G_{1-\theta}$ alkyl and the anyl molely is the above-defined $G_{\Phi/4}$ anyl. Preferably, it is anylalkyl wherein the altry molely thereof is straight orbain alkyl having 1 to 4 carbon atoms and the anyl molely is phenyl. Examples thereof include benzyl, phenetryl, 3-phenyl-

(2089) Tho C₆₋₁₄ any C₁₋₆ alkyl is particularly proferably bonzyl at Re^a and Re^a. (2014) (C₁₋₆ alkyl molety thereof (2070) The C₉₋₁₄ anyl C₁₋₆ alkyloxycarbonyl is anylalkyloxycarbonyl wherein the afkyl molety thereof is at religible to betwo-defined C₉₋₁₄ anyl-C₁₋₆ alkyl. Proferably, life anylalkyloxycarbonyl wherein the afkyl molety thereof include benzyloxycarbonyl, charles thereof include benzyloxycarbonyl.

3

8 phanethyloxycarbony, 3-phorrybropyloxycarbonyl, 2-phorrybropyloxycarbonyl, 4-phorrybropyloxycarbonyl and the like. (d071) The C_{B+L} anyl C_{L_2} altykorycarbonyl is particularly preferably benzyloxycarbonyl at P2. (d071) The cybropyloxycarbonyl as particularly preferably benzyloxycarbonyl at P2. (d071) The cybronally substituted C_{L_2} altyky is the above-defined C_{B+L} altyky. preferably that wherein straight chain or branched chain altyl having 1 to 4 carbon atoms is optionally substituted with 1 to 3 substituent(s), and includes unsubstituted altyl. The substituted(s) is carbon atoms is optionally substituted altyl. The substituted(s), and includes unsubstituted altyl. The substituted(s), altoy, the above-defined C_{B+L} altixynthe, above-defined

2

8

urfluoromethyl or hydroxymethyl et 2, 2° and group D, and methyl at other substituents. [0078] The optionally substituted C₂₋₈ atkenyl is that wherein straight chain or branched chain atkenyl having 2 to 6

ŧ

6

ta(are) selected from the above-defined hatogen atom, hydroxyl group, carboxyl, amino, the above-defined C₁₋₈ alkoxy, the above-defined C₁₋₈ alkoxy, and the above-defined C₁₋₈ alkylamino. Exemples of optionally substituted C₂₋₈ alkylamino. Exemples of optionally substituted C₂₋₈ alkylamino. Exemples of optionally 3-tachexenyl, attempt include vinyl, aftyl. 1-propenyl, teopropenyl, 1-butenyl, 2-butenyl, 1,3-butedenyl, 2-tachexenyl, carbon atoms is optionally substituted by 1 to 3 substituent(s), and includes unsubstituted sikonyl. The substituent(s)

4-methyl-3-pontenyl, 2-carboxylothenyl end the like.

[0079] The optionally substituted G_{2,8} alkonyl is preferably 2-carboxylethenyl at X, and preferably 2-isopentanyl,

3-isobexenyl or 4-methyl-3-penionyl at Fazi.

3-isobexenyl or 4-methyl-3-penionyl at Fazi.

[0077] The optionally substituted G_{2,8} altynyl is that wherein straight chair arroundationed chair altynyl heaving 2 to 6

[0077] The optionally substituted G_{2,8} altynyl is that wherein straight chair uncubalituted altynyl. The substitutent of the carbox desired altynyl, 1-method and the control of the carboxyl, arribo, the above-defined C_{1,8} altynyl in altoxyl group, carboxyl, arribo, the above-defined C_{1,8} altynyl in a straight chair of the carboxyl include othynyl, 1-pro-

ક

ŧ

defined C_{p+q} and is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted anyl. The substituent(s) islant) selected from the above-defined halogen atom, cyano, filtro, the above-defined C_{l+q} slight, the above-defined halogen atom, cyano, filtro, the above-defined C_{l+q} slight, the above-defined C_{l+q} sixanoy, -(CH₂),-CO₁-(CH₂),-CO₁-(CH₂),-CO₂-(CH₂),-CO₂-(CH₂),-CO₃-(CH₂),-CO₄ pynyl, 2-propynyl, 3-butynyl and the like. [2078] The optionally substituted C₂₋₄ altynyl is preferably 2-propynyl at רא²⁰. [2079] The C₆₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from group B is that wherein the above-

8

EP 1 162 188 A1

(0080) Examples therool includo phenyl, naphthyl, anthryl, Indonyl, azutonyl, fluoronyl, phenanthyl, 3-fluorophenyl, 4-fluorophenyl, 3-fluorophenyl, 3-fluorophenyl, 3-fluorophenyl, 3-fluorophenyl, 4-mothyl-phenyl, 3-fluorophenyl, 4-ethorophenyl, 4-ethoro

5

stom or chlorine atom.

(0082) With regard to "C_{6,14} aryl optionably substituted by 1 to 5 substituent(s) selected from group B", it is preferably phenyl, 4-the hoppy at 14-12, R427, Phenyl, 4-the hoppy at 14-12, R427, R429, R

5

[0684] The C₈₊ and planning the substitued by 1 to 8 substituents.
[0684] The C₈₊ and planning substitued by 1 to 8 substituent(s) and includes unabstituent and, the above-defined C₈₊ and period politically substitued by 1 to 8 substituent(s), and includes unabstituent and, The aubstituent(s) is (an).
[0685] Exemples of group D from beluco fluorino store, chore, born, bromine atom, chore, carrier, carrier, and the above-mendional group D (substituents shown under (s) to (s)).
[0685] Exemples of group D from beluco fluorino store, chore, chore, bromine atom, chore, carrier, carbonylinethyl, ethoxycarbonyl, derival, proprise and proprise includes the chore, bromine atom, bromine atom, chore, carbonylinethyl, exemply, t 8

뇮 ä

It is fluorine atom, chlorine stom, mothyt, tort-butyl, carboxyl, mothoxy, carbamoyl, methylihlo, dimothylaminocarbonyl methylauflonyl or acetylamino, most proferably fluorine stom or chlorine stom.

[0090] The heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from group B is that wherein the above-defined hoterocyclic group is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted heterocyclic group. The substituent(s) is clocified in the above-defined heterocyclic group. The substituent(s) is clocified in the above-defined chapter and the above-defined C₁₄ sith, the ab

8

8

à

å

Examples thereof include 2-pyridyl, 3-pyridyl, 4-pyridyl, 3-fluoropyridin-4-yl, 3-chloropyridin-4-yl, 4-chloropy

alkyl and r is 0 or an integer of 1 to 8

ridin-3-yi, pyrazinyi, pyrimidinyi, pyridazinyi, 1,3,6-triazinyi, pyriolyi, pyrazolyi, imidazolyi, 1,2,4-triazolyi, tetrazolyi, zhianyi, tunyi, oxazolyi, z-methythiazol-4-yi, 2,6-dinethythiazol-2-yi, 2,6-dinethythiazol-4-yi, 2,4-dinethythiazol-6-yi, a,4-dinethythiazol-6-yi, a,4-dinethythiazol-6-yi, a,4-dinethythiazol-6-yi, a,4-dinethythiazol-6-yi, a,4-dinethythiazol-6-yi, a,5-dinethythiazol-6-yi, a,5-dinethythiazol-6-4-hydroxypipendino, N-methytoipendin-4-yi, N-(lent-butoxycarbonyl)pipendin-4-yi, N-ecotypipendin-4-yi, N-methytoipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-4-yi, pipendin-yi, -din-din-yi, -din 4-(hydroxymethyl)-piperidino, 2-oxopiperidino, 4-oxopiperidino, 2,2,8,6-tetramethylpiperidino, 2,2,8,6-tetramethyl-

[0932] The histocyclic molety is pretorably a heterocyclic group which is a 5-membered or a 8-membered monopolic group. Examples thereof include pyridyl, pyrazinyl, pyriddinyl, hydrazinyl, 1,3-5-trazinyl, pyrrolyl, pyrrolyl, pyrazolyl, indicazolyl, 1-2,4-flazolyl, tetrazolyl, thenyl, turyl, oxazolyl, isoxazolyl, thizolyl, stotilazolyl, thiadiazolyl, pyrrolldinyl, pperidyl, plecazinyl, norpholinyl, thiamospholinyl and tetrahydropyranyl, and the group Bhere is pretarbly the above defined C-1,4 allyl, the above-defined C-1,4

20

3

ä

prolin-4y.

[googl The heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from group D is that wherein the above-defined heterocyclic group is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted the above-mentioned group D (subheterocyclic group. The substituent(s) is (are) selected from the substituent(s) of the above-mentioned group D (subheterocyclic group. The substituent(s) of the above-mentioned group D (subheterocyclic group.) tituonts shown under (a) to (p)).

to 5 substituent(s) selected from group D. Examples of the group D here include the substituent(s) exemplified for Co.14 any optionally substituted by

8

(2088) Exemples of hoterocyclic group optionally substituted by 1 to 5 substituent(s) selected from group D include 2-pyridyi, 3-pyridyi, 4-pyridyi, 3-duropyridin-4-yi, 3-chloropyridin-4-yi, 4-chloropyridin-3-yi, pyrazinyi, pyrindinyi, pyrazinyi, pyrazinyi, pyrazinyi, indazoyi, indazoy

벊

â pyranyi, quinolyi, isosuinolyi, quinezolinyi, quinoxelyi, phthalezinyi, cinnolinyi, nephthyddinyi, 5,8,7 e-terentydroqui-nolyi, Indolinyi, benzimidezolyi, indolinyi, benzolitanyi, benzolitanyi, benzolitanyi, benzolitanyi and the like. (1987) In addition, the heterocyclic group may be autentituded at the 3-4-4-5-or 6-position of 2-yhidyi, at the 2-4-4-5- or 6-position of 3-pyridyi, at the 2-, 3-, 5- or 6-position of 4-pyridinyi, at the 3-, 4- or 6-position of 2-thienyi, or at the 2-, 4- or 5-position of 3-thlenyf, by fluorine stom, chlorine stom, bromine stom, nitro, methyf, tert-butyf, osrboxyf, trif uoromethyi, hydroxymethyi, methoxymethyi, 2-carboxylothyi, methoxy, carbamoyi, methyithio, dimethylaninocarbo

ηγή, methylisuitionyl or seolylamino.

[0098] At Z and Z, the haterocyclic molety is preferably a heterocyclic group which is a 5-memberod or 8-memberod monocyclic group. Examples thereof include pyridyl, pyractinyl, pyrindidinyl, pyridazinyl, 1,3,5-tinazinyl, pyrindizolyl, 1,3,4-tinazinyl, interest pyrindizolyl, 1,2,4-tinazolyl, interest pyrindizolyl, 1,2,4-tinazolyl, interest pyrindidinyl, pheridyl, piperazinyl, mophelinyl, thienorpholinyl and teterbytropyranyl. The group D here is prefearably the above-defined optionally substituted C_{1,4} elkyl, -(CH₂),-COOR^{4,19}, -(CH₂),-COOR

The C₂₋₈ cyclosityl optionally substituted by 1 to 5 substituent(s) selected from group C is that wherein the defined C₂₋₈ cyclosityl is optionally substituted by the 1 to 5 substituent(s) selected from hydroxyl group, the defined ch₂₋₈ cyclosityl is optionally substituted by the 1 to 5 substituent(s) selected of the hydroxyl group, the defined ch₃₋₈ strong, which may be unaub-

8

EP 1 162 196 A1

2-methylcyclopentyl, 3-methylcyclohezyl, 4-methylcyclohezyl, 4,4-dimethylcyclohezyl, 3,5-dimethylcyclohezyl, 4-tert-bullycyclohezyl, 4-hydroxycyclohezyl, 4-methoxycyclohezyl and 2,3,4,5,6-pentallucorscyclohezyl, 6-tyclohezyl, 4-methoxycyclohezyl, 4-fychohezyl, 4,5,6-pentallucorscyclohezyl, 6-tyclohezyl, 6-ty

group C is preferably cyclopentyf, cyclohexyf, 4-fluorocyclohexyf, 4-methylcyclohexyf, 4,4-dimethylcyclohexyf, 4-tert-butyfcyclohexyf, 4-hydroxycyclohexyf or 4-methoxycyclohexyf, more preferably cyclopentyf or cyclohexyf, particularly

[0104] The C₃₄ cyclosityl optionally substituted by 1 to 5 substituent(s) selected from the above group B is that wherein the above-defined C₃₄ cyclosityl is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted cyclosityl in the above group B. [0108] Specific examples thereof include cyclopropyl, cyclopropyl, cyclopanyl, cyclohayyl, cyclohayyl, cyclopanyl, 3-6-dimethylcyclohayyl, 2-nethylcyclopanyl, 3-methylcyclobayyl, 4-methylcyclopayl, 4-methylcyclopayl,

i

bromine atom, nitro, mathyl, ten-butyl, carboxyl, trifluoromethyl, hydroxymethyl, mathoxymethyl, 2-carboxylethyl, math-oxyl, carbamoyl, mathythilo, dmathytentinocarboxyl, mathyteutionyl or aceylenthol.

[01077] Al cycleshyl motiby, it is preferably cyclopanyl or cyclohasyl, a but C-3, cycloshyl or 4-hydroxycyclohasyl et a 1075 substituent(e) selected from the above group B, it is particularly preferably cyclohasyl or 4-hydroxycyclohasyl et a Pa27 and Pa28

20 [0108] The C_{P-8} cycloalkyl optionally substituted by 1 to 5 substituent(s) selected from group D is that whorein the above-defined C_{P-8} cycloalkyl is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted cycloalkyl. The substitutent(s) is (are) selected from the substitutent(s) of the above-monitoned group D (substituents shown under

(a) to (p)).

[9109] The group D here includes the substituents recited with regard to C₆₋₁₄ aryl optionally substituted by 1 to 5

3

g substituent(s) selected from group D.

(9110) Examples thereof include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclohexyl, 4-fluorocyclohexyl,
2-mathyloyclopenyl, 3-mathyloyclohexyl, 4-mathyloyclohexyl, 4-d-dimethyloyclopenyl,
2-mathyloyclohexyl, 4-hydroxycyclohexyl, 4-mathxycyclohexyl and 2,3,4,5,6-pentafluorocyclohexyl,
(1911) The group D may be, for example, cyclopentyl or cyclohexyl substituted by fluorine atom, chlorine atom,
bromine atom, nâtro, methyl, tort-buryl, carboxyl, trifluoromethyl, hydroxymethyl, methoxymethyl, carboxyletyl, methylsulfonyl or acceptamio.

[1912] The cycloalityl molety is preferably cyclopentyl or cyclohexyl, and at Z and Z, it is particularly preferably

â G [0113] The optionally substituted C_{2.8} cyclosikenyl is that wherein the above-defined C_{2.8} cyclosikenyl is optionally substituted by substituted by substituted by substituted by substituted by substituted from hydroxyl group, the above-defined biogen atom, the above-defined C_{1.8} sikely and the above-defined C_{1.8} sikely, which may be unabstituted. Examples thereof include cyclopropeny, expeciationally, cyclopentenyl, cyclopentenyl, 4-fluoro-2-cyclobasenyl, 4-methyl-2-cyclobasenyl, 4-m

ŝ [0114] The opionally substituted C_{3.2} cyclealizary is particularly preferably cyclohoxenyl at the ring Cy. [0115] The C₆₊₄ eryl C₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from group B is that wherein the above-defined C₆₊₄ eryl C₁₋₆ alkyl is optionally substituted by 1 to 5 substituent(s), and includes unsubstituted anylally. The substituent(s) lated pleated from the above-mentioned group B. [0116] Examples thereof include benzyl, 1-naphthylmethyl, 2-naphthylmethyl, 3-phenytynopyl, 2-phenytynopyl, 4-fachlorobenzyl, 4-fachlorobenz

8 4-methylsutfonylbenzyl, 4-aminosutfonylbenzyl, 3-nitro-4-methoxybenzyl and 4-nitro-3-methoxybenzyl.

(1117) The C_{P-1} anyl C₁₋₄ alkyl ministy is preferably benzyl or phenethyl, particularly preferably benzyl. The group B is preferably the slowe-calined the logen atom, nitro, the above-defined C₁₋₄ alkyl, the above-calined halogen atom, nitro, the above-defined C₁₋₄ alkyl, the above-calined halogen atom, nitro, the above-defined C₁₋₄ alkyl, the above-calined the above-defined C₁₋₄ alkyl, the above-defined C₁₋₄ alkyl alkyl

8 alkyl or (CH₂), OR*1. Examples thorod include fluorine atom, chlorine atom, nitro, methyl, terl-butyl, trifluoromethyl, methoxy or trifluoromethyloxy, particularly preferably fluorine atom or chlorine atom.

[0118] The specific C₆₊₁ any C₁₋₆ alkyl optionally substituted by 1 to 5 substituted(s) selected from group B at Re*1 and R*1 is preferably benzyl, phoneblyl, 3-chlorobenzyl, 4-fart-butybenzyl or 3-trifluoromethylbenzyl, and R*1 is preferably benzyl phoneblyl, 3-chlorobenzyl, 4-trifluoromethylbenzyl, phoneblyl, 3-chlorobenzyl, 4-chlorobenzyl, 3-chlorobenzyl, 4-chlorobenzyl, 4-trifluoromethylbenzyl at R*2°, and 4-chlorobenzyl, 1-trifluoromethylbenzyl at R*2°, and 1-trifluoromethylbenzyl at R*2°, and

[0119] It is penicularly preferably benzyl at other substituents. [0120] The Cq., a sryl Cq., a siryl optionally substituted by 1 to 5 substituent(s) selected from group D is that wherein the above-defined Cq., acryl Cq., attlyd to potionally substituted by 1 to 5 substituent(s), and includes unaubstituted aryl. The substituent(s) is(are) selected from the substituent(s) of the above-mentioned group D (substituents shown under

ymethylleminocarbonyi, hydroxyl group, methoxy, ethoxy, tappropytoxy, hydroxymethyloxy, carboxylmethyloxy, dimbrytambocarbonylmethyloxy, ambo, methylambo, diethylemino, dethylemino, ceeplyamino, methylambo, diethylemino, diethylemino, dethylemino, methylambosulionyl, notypedialinyl, embrytambosulionyl, embrytambyl, embrytambosulionyl, embrytambosulionyl, embrytambyl, embrytambosulionyl, embryt methoxybenzyi, 4-carbamoybenzyi, 4-methylthlobenzyi, 4-(dimethylaminocarbonyi)benzyi, 4-methylaultonybenzyi, 4-(acetylamino)benzyi, 4-cyanobenzyi, 4-acetylbenzyi, 4-aminobenzyi, 4-dimethylaminobenzyi, 4-(methylaultonylamino) benzyi, 4-methylaultinybenzyi, 4-aminosultonybenzyi, (3-nitro-4-methoxyphenyi)methyl and (4-nitro-3-methoxyphenyi)methyl and (4nyi, isopropylaminocarbonyi, dimethylaminocarbonyi, diethylaminocarbonyi, (2-hydroxyethyi)aminocarbonyi, (carbox isopropyl, tert-butyl, trifluoromethyl, hydroxymethyl, 2-hydroxyethyl, methoxymethyl, 2-carboxylethyl, methoxycarbo-nylmethyl, ethoxycarbonylmethyl, ecetyl, carboxyl, methoxycarbonyl, ethoxycarbonyl, carbamoyl, methylaminocarbo-(e) to (p)). [0121] Examplos of group D include fluorine siom, chlorine stom, bromine stom, nitro, cysne, methyl, ethyl, propyl,

[013] At Zend Z, tho C₆₋₁₄ eryl C₁₋₄ alkyl molety is prole rably bonzyl or phonethyl, and the group D here is preferably the above-defined hetegon etcm, titre, the above-defined optionally substituted C₁₋₄ alkyl, -(CH₃),-COOR*** (-(CH₃),-COOR***), -(CH₃),-COOR*** (-(CH₃),-COOR***), -(CH₃),-COOR*** (-(CH₃),-COOR***), -(CH₃),-COOR***, -(CH₃),-COOR***, -(CH₃),-COOR***, -(CH₃),-COOR***, -(CH₃),-COOR***, -(CH₃),-COOR***, -(CH₃),-COOR**, -(CH₃),-CO

[0125] It is particularly preferably the above-defined chalges atom, the above-defined optionally substituted C₁₋₈ alkyl, C(CH₂)-COCR¹²⁺, -(CH₂)-CONR²⁷Re²², -(CH₂)-COR¹²⁺ or - (CH₂)-S(O)₂-Re²², Examples thereof include florence atom, cholorine atom

1,2,4-frezolyf, telrezolyf, thienyf, furyf, oxezolyf, isoxezolyf, thiazolyf, isothlezolyf, thiadiazolyf, pyrrolldinyf, piperidyf, piperidyf, propholinyf, thornopholinyf, thornopholinyf, and the allyf modely thereof is preferably straight chain allyf having 1 to 4 carbon atoms. The group B here is preferably the above-defined halogen atom, the above-defined talkyf, the above-defined C_{1,4} allyf, the above-defined halogenatind C_{1,4} allyf, the above-defined C_{1,4} allyf, the above-defined C_{1,4} allyf, the above-defined the Conference of Conf group. Examples thereef include pyridyl, pyrazinyl, pyrimidinyl, pyridazinyl, 1,3,6-triazinyl, pyrrolyl, pyrazelyl, imidazelyl

[0129] Examples of heterocycle C₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from group B preferably neutron and the Apyridymentyl, 3-pyridymentyl, 3-chicropyridin-4-ymentyl, 1-pyridymentyl, piperidin-4-ymentyl, 1-menylpiperidin-4-ymentyl, 1-de-hydroxyngrowyl) piperidin-4-ymentyl, 1-de-hydroxyngrowyl piperidin-4-ymentyl, 1-de-hydroxyngrowy

8

EP 1 162 198 A1

neihyl, 2,4-dimatrylithazolin-6-yimothyl and 4-mothylihazol-2-yimothyl. Particularly protonathy, it is 2-pyridymathyl, 2-dimatrylithazolin-6-yimothyl, 4-pyridymathyl, 2-dimatrylithazolin-4-yimothyl, 2-pyridymathyl, 2-mothylipoladin-4-yimothyl, 2-qyridymathyl, pipardin-4-yimothyl, 1-mothylipoladin-4-yimothyl, 2-querylipoladin-4-yimothyl, 1-querylipoladin-4-yimothyl, 1-querylipoladin-4-yimothyl, 2-querylipoladin-4-yimothyl, 2-querylipoladin-4-yimothyl, 2-querylipoladin-4-yimothyl at Paza and Paza, and 4-pyridymathyl or 4-mothylithazol-2-yimothyl at Paza and Paza. [0130] The Cyglosithyl Cy₄₋alixyl optionatly substituted by 1 to 5 substituent(s), and includes a that whosein the above-defined Cyglosithyl Cy₄₋alixyl soptionatly substituted by 1 to 5 substituent(s), and includes a that whosein the above-defined Cyglosithylipolyl. The substituents are selected from the above group B.

ō [0131] Specific examples thereof include cyclopropylmathyl, cyclobunylmathyl, cyclopentylmathyl, cyclopentyl

ciohexytriethyi.

[[0132] Asso exempillied are those wherein cyclopenlylmethyl or cyclohexytmethyl is substituted by fluorine atom, chlorine atom, chlorine

3

wherein each symbol is as defined above.

[0135] G1, G2, G3 and G5 are each preferably (C-R1), (C-R2), (C-R2) and (C-R2), G3 is preferably a nitrogen atom, and G5, G3 and G6 are preferably a carbon atom. G7 is preferably C(-R2) or unsubstituted nitrogen atom, wherein R7 is preferably hydrogen atom.

[0136] A preferable corribination is G3 of (C-R2) and G5 of a carbon atom, particularly preferably G3 of (C-R2), G5 of a carbon atom and G5 of a nitrogen atom, most preferably G2 of (C-R2), G5 of a carbon atom, G3 of a nitrogen atom and G2 of unsubstituted nitrogen atom.

is(are) preferably a nitrogen atom, specifically preferably

ĸ

more preferably

EP 1 162 196 A1

most proforably

[0138] R1 and R4 are preferably hydrogen atom, R2 is preferably carboxyt, -COOR±1, -COOR±2R2 or -SO₂Re2 (each symbol is as defined above), particularly preferably carboxyt, -COOR±1 or -SO₂Re2, more preferably carboxyt or -COOR±1, most preferably carboxyt or -COOR±1, most preferably carboxyt. A3 is preferably hydrogen atom or -OR±6 (R±6 is as defined above), particularly

[6139] The ring Cy and ring Cy are preferably cyclopentyl, cyclohexyl, cyclohexyl or tetrahydrothlopyranyl, particularly preferably cyclopentyl, cyclohexyl or tetrahydrothlopyranyl, particularly preferably cyclopentyl, cyclohoxyl or cyclohoxyl, cyclohoxyl

or pyridayl, and most pretramaby phonyl.

[0141] The ring B and ring B are pretramably C_{1,4} anyl or hoterocyclic group, specifically preferably, phenyl, pyridyl, prezinyl, pyridalinyl, 1,3 6-trazinyl, pyridyl, pyridyl, pyridyl, pyridyl, pyridyl, pyridalinyl, 1,3 6-trazinyl, pyridyl, pyridyl or bhazolyl, and most perfereably phonyl, pyridyl, pyridyl, pyridyl or bhazolyl, and most perfereably phonyl, pyridyl, pyridyl, pyridyl, pyridyl, pyridyl, and most perfereably phonyl, and most perfereably pyridylen atoms, when ring A is phonyl, Pi and Pi preferably are present at an ortho position from 96. The same applies to Pi and Pi.

[0143] Y is preferably, (CH₂)_m, or (CH₂)_m, APP-12-CH₂-15, CONH-CHR-14, - (CH₂)_m, APP-12-CH₂-15, - CONH-CHR-14, - (CH₂)_m, APP-12-CH₂-15, - CONH-CHR-14, - (CH₂)_m, - CONH-CH₂-16, - (CH₂)_m, - CPR-13-P(-14)_m, - (CH₂)_m, - (CH₂)_m

$$-\left(\mathrm{CH}_{2}\right)_{a}$$
 $+\left(\mathrm{B}\right)$ $-\left(\mathrm{Z}^{\prime}\right)_{\mathbf{A}^{\prime}}$

molety is proforably symmetric. The preferable mode of n, ring B, Z and w and the preferable mode of n', ring B', Z and w and the preferable mode of n', ring B', Z and w and the preferable mode of n', ring B', Z and w are the same.

[0145] Whon ring A is phenyl, X or Y is preferably present at the pera-position relative to G⁰. When ring B and ring B' are phenyl, Z is preferably present at the ortho or meta-position relative to X', it is preferable that the 3-position on phenyl each have one substituent.

[0147] Whon ring B is intactly, Y is preferably substituted at the 5-position, and Z is preferably substituted at the 5-position, and Z is preferably substituted at the 5-position, and Z is preferably substituted at the 5-position.

substituted at the 5-position, and Z is preferably substituted at the 2-position, the 4-position or the 2-position and the

[0148] Z and Z are preferably group D, "C₉₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from group D" or "netercoyclic group optionally substituted by 1 to 5 substituent(s) selected from group D", perfectlerly preferably

g

t group D or C_{6,14} anyl optionally substituted by 1 to 5 substituent(s) selected from group D*.

[1949] More preferably, they are the above-defined helogen atom, nitro, the above-defined optionally substituted C_{1,4} slipy, 4CH₂h₂COO(Pari²), CON₂h₃COO(Pari²), CON₂h₃COO(Pari²), CON₄Pari²CO₁Ch₄h₃COO(Pari²), CON₄Pari²CO₁Ch₄h₃COO(Pari²), CON₄Pari²CO₁Ch₄h₃COO(Pari²), CON₄Pari²CO₁Ch₄h₃COO(Pari²), CON₄Pari²CO₁Ch₄h₃COO(Pari²), CON₄Pari²CO₁Ch₄ slipy or heterocyclic group optionally substituted by these.

With regard to Z and Z', the preferable mode of group D that identify state of Ch₄ slipy or heterocyclic group are the proferable mode of group D that substitutes of C₁, at 9N, 2₂ cyclosityl, C₆ statiff or heterocyclic group are the same, whorigh they may be the same with or different from each other.

[1950] Specific examples of the substitutent preferably include fluorine stom, chlorine stom, bromine stom, premony, stome of the substituted preferably include fluorine stom, ethotoxymoliny, 2-curboxylethy, methoxycarbony, ethoxycarbony, ethoxyca

EP 1 182 198 A1

3 3-methoxybenzóylamino, 3-pyrárylcatbonylamino, 4-methyphenylautionylamino, 2-iniazolylaminosutonyl, 2-pyrigylaminosutonyl, benzylaminocatbonyl, N-benzyl-N-methytaminocatbonyl, 4-pyridymethyl-aminocatbonyl, 3-hydroxypilodohoxylmethylaminocatbonyl, 2-hydroxypiloyloxy, 3-hydroxyproplycay, 2-byrdoxyproplycay, 3-hydroxypilodinocatbonyl, 3-d-dihydroxypiperidinocatbonyl, 4-methyzminocatbonyl, 2-z, 8-tetramethyl-byridypiperidinocatbonyl, 4-methypiperidinocatbonyl, 2-z, 8-tetramethyl-byridypiperidinocatbonyl, 4-methypiperidinocatbonyl, 1-d-actionathylaminocatbonyl, 1-d-act rahydropyranyloxy, 2-pyrdylmathyloxy, 3-pyrldylmathyloxy, 2-chloropyrddin-4-ylmothyloxy, 4-pyrldylmathyloxy, 2-pbridylmathyloxy, 3-plperdylmathyloxy, 4-plperdylmathyloxy, 1-mathyloperddin-4-ylmathyloxy, 1-acetyloperddin-4-ylmathyloxy, 1-acetyloperddin-4-ylmathyloxy, 1-acetyloperddin-4-ylmathyloxy, 1-acetyloperddin-4-ylmathyloxy, 2-mathynthazolin-4-yloxy, 2-4-dimothythhazolin-5-yloxy, dimothylamlocetaropylmathyloxy, piperddin-carbonylmathyloxy, 2-mathynthazol-4-yl, (2-methylthiazol-4-yl) methyloxy, (2-4-dimothydhazol-6-yl)methyloxy, benzoyl, 3-fuorobenzoyl, 4-chlobonzylamino, 3-f-dichlorobonzylamino, 4-fulluoromathylamazylamino, 2-pyrldylmethylamino, benzoylamino, 4-chlobonzylamino, 3-f-dichlorobonzylamino, 4-fulluoromathylamaylamino, 2-pyrldylmethylamino, benzoylamino, 4-chloyloxy and 4-methythlazol-2-yimethyloxy. robenzoylamino, 4-trilluoromethylbenzoylamino, 3,5-dichlorobenzoylamino, 3-nitro-4-methoxybenzoylamino, 4-nitro

(1811) Particularly professibe szampiles of the substituent include fluorine atom, chlorine atom, bromine atom, nitro populo, melhyl, hydroxymelhyl, carbanyl, carbanyl, methydaminocarbonyl, telepoxymethyl-aminocarbonyl, dimethydaminocarbonyl, (lethydaminocarbonyl, clarbonyl), carbanyl, 3

ô

pound.
[0153] The compounds of the above-mentioned formula [i] or [ii] have various isomers. For example, E compound and Z compound are present as geometric isomers, and when the compound has an asymmetric curbon, an ensuturner and a disservemer are present due to the asymmetric curbon. A laudomor may be also present. The present invention encompasses all of these isomers and mixtures thereof.

8

8 [0155] The present invention also oncompasses prodrug and matabolite of each compound.

(0157] A prodrug means a derivative of the compound of the present invention, which is capable of chemical or metabolic decomposition, which a howe inherent efficacy by revering to the original compound effer administration to a body, and which includes saits and complexes without a covalent bond.

(0158) When the inventive compound is used as a phermacoutical preparation, the inventive compound is generally

-pyridyl, 4-pyridyl, 2-pyrimidinyl, 5-tetrazolyl, piporidino, piperidinocarbonyl, 4-hydroxypiperidinocarbonyl, 1-piperazi-

E

8

Ġ

â

admixed with pharmaceutically accopable carriers, excipients, ciliuents, binders, delantegrators, atabilizers, presorvatives, butlers, emutafilers, eromatics, coloring agents, exvesioners, thetesnest, correctives, soutbilizers, and other at-diffives such as water, regotable oil, alcohol such as ethanol, benzyl alcohol and the like, polydithylene gives, giyeerol ritacestate, gelatin, factoes, carbohydratie such as starch and the like, magnetium stearate, tate, lanolin, patrostum and the like, and prepared into a decage form of tablets, pilit, powders, granules, suppositoria, preciona, eye dropt the like, and prepared into a decage form of tablets, pilit, powders, granules, suppositoria, manualions, syrups and the like, which can be administered inducts, capsules, troches, serosois, olidits, suspensions, amulaions, syrups and the like, which can be administered systemically or topically and orally or parenterally.

(0155) While the dose varies depending on the age, body weight, general condition, treatment effect, administration route and the lite, it is from 0.1 mg to 1 g for an adult per dose, which is given one to several times a day.

(0160) The prophylaxis of hepatitis C means, for example, administration of a pharmacuical agent to an individual round to carry an KCV by a test and the like but without a symptom of hepatitis C, or to an individual who shows an improved disease state of hepatitis after a treatment of hepatitis C, but who still carries an HCV and its associated with a risk of recurrence of hepatitis.

[0161] Examples of the production method of the compound to be used for the practice of the present invention are given in the following. However, the production method of the compound of the present invention is not limited to those

(0182) Even it no directly corresponding disclosure is found in the following Production Mothods, the steps may be modified for efficient production of the compound, such as introduction of a protecting group into a functional group with deprotection in a subsequent step, and changing the order of Production Methods and steps.

[0183] The treatment after reaction in each step may be conventional once, for which typical methods, such as isolation and purification, crystalization, recrystalization, elics gel chromatography, preparative HPLC and the like, can be appropriately selected and combined.

Production Method 1

[0184] In this Production Method, a banzimidazole compound is formed from a nitrobanzane compound

[0185]

wherein Hal is halogen atom, such as chibrine atom, bromine atom and the like, Ref is halogen atom, such as chlorine atom, bromine atom and the like, or hydroxyl group, and other symbols are as defined above.

EP 1 162 196 A1

Stop 1

(0188) A compound [1] obtained by a conventional method or a commercially evallable compound [1] is reacted with affine compound [2] in a solvent such as N.N-dimathytionnamide (DNF), acetonitite, teirahytionnamide (DNF) acetonitite, teirahytionnamide and the like in the presence or absence of a base such as potassium actionate, triethytamine, potassium t-buloade and the like at room temperature or with hosting to give compound [3].

[0187] The compound [3] is hydrogenated in a solvent such as methanol, ethanol, THF, ethyl acetate, acatic acid, water and the like in the presence of a catalyst such as patiadium carbon, patiadium hydroxide, plathrum oxide, flancy nickal and the like at room temperature or with heating to give compound [4]. In addition, compound [3] is reduced with a reducing agent such as zinc, iron, lin(i) chiefde, sodium suffice and the like, or reacted with hydrazine in the presence of tren(iii) chiefde to give compound [4].

This chloroform, eitry acelete, methylene chlorde, lothene and the like using a condensing agent such as disyclohestylechoodlinide, i. Isruhi-4-(3 dimethylaminopropi)canbodlinide hydrochloride, diphorylphosphoryl azide and the like
and, where necessary, adding N. hydroxysucchimide, i. hydroxybonzoritazole and the like to give amide compound
(6), Altornatively, amide compound (6) can be obtained from compound (6) as flower. The acetoxylic acid compound
(6) is converted to an acid halide derived with bioloyyl chloride, pashyl chioride and the like, or an acide cester (6.5,
mixed socid anhydride derived with eithyl chiorocerbonate and the like), which is then reacted in the prosence of a base, such as triethylamine, potassium carbonate, pyridine and the like, or in an amine solvent, such as pyridine and the like, to give amide compound [6]. [0168] The compound [4] is condensed with carboxylic acid compound [5] In a solvent such as DMF, acetonitrile

(0169) The compound [6] is heated in a solvent such as ethenol, methanol, totuene, DMF, chloroform and the like or without a solvent in the presence of an acid such as accrite acid, formize acid, hydrochloric acid, dilute suffurite acid, phosphoric acid, orbyphosphorio acid, p-toluenesuiforic acid and the like, a habgenating agent such as zinc chloride, phosphorus oxychloride, thenyl chloride and the like or acid enhydride such as acetic anhydride and the like, to allow cyclization to give compound [1-2].

Production Method 1-2

[0170] This Production Mathod is an alternative method for producing compound (1-2).

ŝ

wherein each symbol is as defined above

Step 1

[0171] The compound [3] obtained in the same manner as in Step 1 of Production Method 1-1 is subjected to amide condensation with compound [5] in the same manner as in Step 3 of Production Method 1-1 to give compound [7].

Stop 2

ö [0172] The compound [7] is reduced in the same manner as in Step 2 of Production Mathod 1-1 to give compound [8].

Step 3

75 [0173] The compound [8] is subjected to cyclization in the same manner as in Step 4 of Production Method 1-1 to give compound [1-2].

Production Method 1-3

[0174]

8

ŝ ĸ Ξ

8 wherein R²² is alkyl such as methyl, ethyl and the like, and other symbols are as defined above.

[9175] The compound [4] is reacted with initiate compound [9] in a selvent such as mothanol, obtained acid, DMF, THF, chloroform and the like at from temperature or with heating to give compound [1-2].

[0176] In actilitin, compound [4] may be reacted with attenyed compound [10] in a solvent such as aceita acid, formic acid, aceitahithe, DMF, nitroborzene, toluene and the like in the presence or absence of an oxidizing agent such as benzefurexan, manganese disside, 2,3-dishlero-5,6-disyno-p-banzbquinone, lodine, potassium ferriesprance and the like in the terminative potassium ferriesprance disside, 2,3-dishlero-5,6-disyno-p-banzbquinone, lodine, potassium ferriesprance and the like with heating to give compound [1-2].

of polyphosphoric acid, phosphoric acid, phosphorus exychloride, hydrochloric acid and the like to give compound [1-2]

EP 1 162 188 A1

Production Method 2

is applicable irrespective of the position of substitution. [0178] In this Production Method, conversion of the substituents (R1, R2, R4) on the benzene ring of benzimidszele is shown. While a method of converting R2 when R1, R3 and R4 are hydrogen atoms is shown, this Production Method

Production Method 2-1

[0179] Conversion of carboxytic acid ester molety to amide

wherein E is a single bond, $\cdot(CH_2)_*$, $\cdot O\cdot(CH_3)_*$, $\cdot O\cdot(CH_3)_*$ (wherein s is an integer of 1 to 6), Ref. Ref and Ref are C_{1+0} sloy), and other symbols are as defined above.

Step 1

[0180] The compound [1-2-1] obtained in the same manner as in the above-mentioned Production Method is subjected to hydrolysis in a selvent such as methanol, ethanol, THF, dozane and the like, or in a mixed selvent of these solvents and water under basic conditions with sediem hydroidele, potestium products, potestium enhancers, lithium hydroides and water under basic conditions with addism hydroided potestium hydroide, potestium cancers, and and the like or under solds conditions with hydrochloric sold, sulfurb sold and the like to give compound [1-2-2], and the like or under solds conditions with hydrochloric sold, sulfurb sold and the like to give compound [1-2-2].

낞 [0181] The compound [1-2-2] is reacted with compound [12] in the same manner as in Step 3 of Production Method 1-1 to give compound [1-2-3].

[0182] Conversion of cyano group to substituted amidine group

Production Method 2-2

8 wherein each symbol is as defined above.

[0183] The compound [1-24] obtained in the asme manner as in the above-mentioned Production Method is reacted [0183]. The compound [1-2-5] with hydroxylamine in a solvent such as water, methanel, athanel, THF, DMF and the like to give compound [1-2-5]. When a sait of hydroxylamine such as hydroxholds and the like is used, the reaction is carried out in the presence of a base such as sodium hydrogencerbonete, sodium hydroxide, triethylamine and the like.

Production Method 2-3

[0184] Conversion of suffonic acid ester molety to suffonic acid

wherein Rst is C₁₋₆ alky4, and other symbols are as defined above.

(1985) The compound [1-2-6] bitsined in the same manner as in the above-mentioned Production Method is reacted with bodide ant such as sodium iodide, lithium iodide and the like, bromide salt such as sodium bromide, trimsihyfammonium bromide and the like, armine such as pyridine, trimsihyfamine, triazole and the like, phosphine such as triphenylphosphine and the like in a solvent such as DMF, dimethyf suffoxide (DMSO), acctonitrile, methenol, ethanol, water and the like with heating to give compound (j-2-7).

Production Method 3

[0186] This Production Method relates to convention of the substituent(s) on phenyl group at the 2-position of ben-zimidazole. This Production Method can be used even when phonyl is a different ring.

Production Method 3-1

[0187] Conversion of hydroxyl group to ether

wherein R^{a7} is optionally substituted alkyl corresponding to R^{a11} , G^1 is a single bond, *- $(CH_2)_n$ -, *- $(CH_2)_n$ -Q-, *- $(CH_2)_n$ -Q-, *- $(CH_2)_n$ -, wherein * show the side to be bended to R^{a1} , and other symbols are as defined

(0188) When FY of compound (13) is halogen atom, compound (1-2-8) obtained in the same manner as in the above-monitioned Production Method is reacted with compound (13) in a solvent such as DMF, DMSO, acetonitrie, ethanol, THF and the like in the presence of a base such as sodium hydride, sodium hydroxide, potassium hydroxide, potassium.

ô

EP 1 162 196 A1

carbonate, sodium ethoxide, potassium t-butoxide and the like at room temperature or with heating to give compound

[H-2-1] When R⁻¹ of compound [13] is hydroxyl group, the hydroxyl group of compound [13] is converted to halogen group with thinnyl chloride, phosphona tribromide, carbon (strabnomide-triphenylphosphine and the like and reacted with compound [12-8] by the attermantioned method to give compound [112-1], in this case, compound [12-9] may be with compound [12-9] to reaction with compound [13] in a solvent such a IS-NF, accionizitie, THF and the like using triphenylphosphine - distriyl accidearboxylate and the like to give compound [12-1].

[0198] The compound [12-9] can be obtained in the same manner from compound [12-9] and compound [14].

Production Method 3-2

[0191] Conversion of nitro to substituted amino group

15

$$R^{1} \longrightarrow R^{1} \longrightarrow$$

\$ wherein R^{cl} is $C_{1:0}$ sity), G^2 is "-(CH₂)," or "-CHR*15, G^2 is -CO-, "-CO_{2"}, "-CONH- or -SO_{2"}, and other symbols are as defined above.

Step 1

8 (0192) The nitro compound (i-2-10) obtained in the same manner as in the above-mentioned Production Method is reacted in the same manner as in Step 2 of Production Method 1-1 to give compound (I-2-11).

8 [0193] The compound [+2-11] is alkylated with compound [15] in the same manner as in Production Method 3-1 to give compound [1-2-2].

(0184) When G3 of compound [16] is -CO-, -CO₂- or -CONH-, compound [1-2-11] is acylated with compound [16] in the same manner as in Step 3 of Production Method 1-1 to give compound [1-2-3].
[3185] When G3 of compound [16] is -SO₂- suffortyfation is conducted using sufferyl halide instead of acid halide used in Step 3 of Production Method 1-1 to give compound [11-2-3].

[0198] The compound (F2-11) is acylated with compound [17] in the same manner as above to give compound (F-2-12).

õ [0197] This Production Method is applied in the same menner as above to give disubstituted compounds (tertiary arthre) of compound (1i-2-2), compound (1i-2-3) and compound (1-2-12).

Production Method 3-3

[0198] Conversion of carboxytic acid eater molety to amide

wherein \mathbb{R}^{a_0} is G_{+a} sity(, \mathbb{G}^a is #-($\mathbb{G}H_2$)_{*}, #-($\mathbb{G}H_2$)₀-NH- or #-CHR¹¹⁴-wherein # shows the side that is bounded to smine and other symbols are as defined above.

(0199) The compound [1-2-13] obtained in the same manner as in the above-mentioned Production Method is reacted in the same manner as in Step 1 of Production Method 2-1 to give compound [1-2-14].

Step 2

Ĝ

[0200] The compound [I-2-14] is reacted with compound [18] in the same manner as in Step 2 of Production Method 2-1 to give compound (II-2-4).

[0201] The compound [I-2-15] is obtained from compound [I-2-14] and compound [19] in the same manner as above.

Production Method 4

૪

(2202) In this Production Method, additional substituent(s) is(are) introduced into ring B on phenyl group that substitutes the 2-position of benzimidazole. This Production Method is applicable even when phenyl is a different ring.

Production Method 4-1

2

[0203] Direct bonding of ring 2" to ring B

EP 1 162 198 A1

wherein ting 2"-M is anyl metal compound, ring 2" motely is optionally substituted C₈₋₁₄ anyl or optionally substituted heteropycitic group corresponding to substituent 2, and the metal motely contains borns, zinc, tin, magnestirm and the like, such as phenylbornnic action, or is 0, 1 or 2, and other symbols are as defined above.

[02304] The compound [II-2-5] obtained in the same manner as in the above-mentioned Production Mathod is reacted

[1,3-bis(diphenylphosphino)-propane)nickei(ii) chloride and the like, and a base such as potassium carbonate, potassium hydrogencarbonate, sodium hydrogencarbonate, potassium phosphate, tristhylamine and the like at room temperature or with heating, to give compound [11-2-6]. with anyl metal compound [20] in a solvent such as DMF, acotontrile, 1,2-dimethoxyethane, TMF, toltene, water and the like in the presence of a patiedum catalyst such as totrakta(triphonytphosphino)-patiedum, bis(triphonytphosphino) patiedum catalyst such as rickel chiorida, patiedum(1) dichloride, patiedum acotate - triphenytphosphine and the like, a nickel catalyst such as nickel chiorida,

Production Method 4-2

[0205] Conversion of hydroxyl group to ether

wherein R¹⁰ is -R²⁰ or -(CH₂)_p-COR²⁰ corresponding to substituent Z, and other symbots are as defined abovo. (0206) The compound [II-2-7] obtained in the same manner as in the above-mentioned Production Method is reaged with compound [21] in the same manner as in Production Method 3-1 to give compound [II-2-8].

Production Method 4-3

[0207] Synthesis in advance of ring B part such as compound [13] in Production Mathod 3-1

wherein Re¹¹ is leaving group such as bromine atom, lodino atom, trifluoromethanesulfonyloxy and the like, Re¹⁸ is formyl, carboxyl or carboxylic acid ester such as methoxycarbonyl, othoxycarbonyl, ten-butoxycarbonyl and the like, and other symbols are as defined above.

25

[0208] Commercially evaliable compound [22] or compound [22] obtained by a conventional mathod is reacted with any metal compound [20] in the same manner as in Production Mothod 4-1 to give compound [23].

Step 2

(0211) The compound [24] obtained in the same manner as in the above-mentioned Production Mathod is reacted in a solvent such as 1,4-dioxane, distryl ether, 11F, dichibromethane, chlorotorm, toluene and the like with a halogentating agent, such as phosphorate percendular, phosphorate tribromide, thionyl chloride and the like, in the presence of a tentiary smike such as phyridine and the like to give compound [25]. (0209) The compound [23] obtained in the same manner as in the above-mentioned Production Method is reduced according to a conventional method to give compound [24]. The conventional method to give compound [24] is reacted within a solvent such as methanol, shanol, THF and the like in the [0210]. For example, compound [23] is reacted within a solvent such as methanol, and the like under cooling to presence of a reducing agent such as ithium abunihum hydride, sodium borehydride and the like under cooling to Step 3 heating to give compound [24].

Ġ

[0212] The compound [24] or [25] obtained in the same manner as in the above-monitoned Production Method is reacted with compound [1-2-8] in the same manner as in Production Method 3-1 to give compound [11-2-8].

8

EP 1 162 198 A1

Production Method 4-4

[0213]

$$(2)_{w} \underbrace{\begin{pmatrix} B \\ B \end{pmatrix}}_{\text{Hal}} \underbrace{\begin{pmatrix} \text{Step 1} \\ \text{Z} \end{pmatrix}_{w}}_{\text{[42]}} \underbrace{\begin{pmatrix} Z' \end{pmatrix}_{w'} - \begin{pmatrix} B' \\ B' \end{pmatrix}}_{\text{[43]}} - \frac{G'}{\text{CHO}}}_{\text{[43]}}$$

$$(2)_{w} \underbrace{\begin{pmatrix} B \\ B' \end{pmatrix}}_{\text{[5]}} \underbrace{\begin{pmatrix} Z' \end{pmatrix}_{w'}}_{\text{[45]}} + \frac{G'}{\text{CHO}}}_{\text{[45]}}$$

$$(2)_{w} \underbrace{\begin{pmatrix} B \\ B' \end{pmatrix}}_{\text{[6]}} \underbrace{\begin{pmatrix} Z' \\ Z' \end{pmatrix}_{w'}}_{\text{[6]}} + \frac{G'}{\text{CHO}}}_{\text{[45]}}$$

wherein M' is a metal auch as magnesium, lithium, zinc and the like, and other symbols are as defined above.

8

[0214] Commercially available compound [41] or compound [41] obtained by a conventional mothod is converted to any metal reagont by a conventional method to give compound [41].

[0215] For example, when M' is magnesium, magnesium is neared with compound [41] in a selvent such as THF, idelity ether, benzene, reluene and the like, preferably THF, from cooling to heating preferably at -100°C to give compound [42].

ä

(0216) The compound (42) obtained in the same manner as in the above-mentioned Production Method is readed with compound (43) to give compound (44), whith compound (43) to give compound (44).

[0217] The compound (42) to exceled in a solvent such as diethyl ether, benzene, toluene, THF and the like, preferably THF, from cooling to room temperature, preferably at -100°C to 30°C to give compound (44).

[0218] The compound [44] obtained in the same manner as in the above-mentioned Production Method is halogen-ated in the same manner as in Step 3 of Production Method 4-3 to give compound [45].

[0219] The compound [44] is reacted with thionylichionds and pyridine preferably intoluene solvent to give compound

[0220] When compound [45] is symmetric, namely, when the ring B-(Z)w molety and the ring B-(Z)w molety are the same, compound [42] is reacted with formate such as methyl formate, ethyl formate and the like, preferably ethyl formate, in a solvent such as a distribly ether, beraces, closers. THF and the like, preferably THF, from cooling to room temperature, preferably at -100°C to 30°C, to give compound [45].

Production Method 4-5

8

[0221] Mothod including steps to introduce a protecting group into a functional group

\$

wherein Re¹³ is carboxylic acid protecting group such as ten-buryl and the like, Re¹⁴ is carboxylic acid protecting group such as mathy and the like and other symbols are as defined above. Step 1 [0222] Commercially available compound [26] or compound [28] obtained by a conventional method is protected by a conventional method to give compound [27].

[0223] For example, when Re¹³ is ten-buryl, compound [26] is convented to acid halido with thionyl chloride, oxalyl colorida and the like in a selvent such as THF, chloroform, dichloromethane, toluene and the like, and reacted with chlorida and the like.

potassium ten-butoxide to give compound [27].

[0224] As used herein, Re13 may be a different protecting group as long as it is not removed during the Step 2 or Step 3 but removed in Step 4 without affecting -CO₂-Re14.

Step 2

Ġ [0225] The methyl group of compound [27] obtained in the same manner as in the above-mentioned Production Method is convented to bramomethyl with N-bromosucchimide and N,N-ezobisisoburyronitrile and reacted with compound [1-2-16] in the same manner as in Production Method 3-1 to give compound [1-2-10].

Step 3

[0226] The compound [I+2-10] obtained in the same manner as in the above-mentioned Production Method is reacted with anyt metal compound [20] in the same manner as in Production Method 4-1 to give compound [I+2-11].

8

(0227) The R^{e13} of the compound (II-2-11) obtained in the same manner as in the above-mentioned Production Methods is removed by a conventional method to give compound (II-2-12).

[0228] The presenting group of carboxylic acid can be removed by a conventional depretaction method according to the protecting group. In this Step, the conditions free from reaction of R^{e14} are preferable. For example, when R^{e13} is

EP 1 162 198 A1

torf-butyl, compound [II-2-11] is treated with trifluoreacelio acid in a solvent such as dichloremethane, chlorolorm and the like to give compound [II-2-12].

compound [II-2-13]. [0229] The compound [1:2:12] obtained in the same manner as in the above-mentioned Production Method is sub-jected to artifa condensation with compound [28] in the same manner as in Step 3 of Production Method 1-1 to give

ŏ Step 6

ĕ

[0230] The compound (II-2-13) obtained in the same manner as in the above-mentioned Production Method is de-protected in the same manner as in Step 1 of Production Method 2-1 to give compound [II-2-14]. [0231] As used herein, Pri's is preferably a protecting group that does not react during the Step 1 through Step 5 but removed in this Step.

[0232] For example, when R⁴⁴ is methyl, compound [11-2-13] is reacted in an abohol solvent such as methanol, ethanol, r-propanol, indeprepanol and the like or a mixed solvent of abond solvent and water in the presence of a base such as petassium exforated, and exiting responsive and the like from cooling to heating for deprotection, followed by additying the reaction solution to give compound [11-2-14].

Production Method 6

[0233] Formation of Indole ring

25
$$H_{2} | -\frac{A}{A} + \frac{B}{B} + (2)e^{-\frac{1}{2}} + \frac{e^{4/4} - c = c - A}{R^{4/4} - c = c - A} + \frac{B}{B} + (2)e^{-\frac{1}{2}}$$
25
$$I(20) | I(20) | I(20)$$

wherein R^{ets} is protecting group such as trimethylsily, tenbutyldimethylsily, ten-butyldiphenylsilyl and the like, and other symbols are as defined above.

Step 1

8

(234) The compound (29) obtained in the same manner as in the above-mentioned Production Method or conventional method is reacted with compound (30) in a solvent such as DMF, accolamite, 1.2 dismolocyprithers, THF, totarene, water and the like using a palledium catalyst such as tearbidictiphenylphosphina) better by better by palledium catalyst such as tearbidictiphenylphosphina palledium, betterphosphips catalyst such as tearbidium and the like, as copper catalyst such as copperfy locidies and the like or a meture thereof, and in the presence of a beas such as poinsatium carbonate, potassium tydrogen-carbonate, sodium hydrogen-carbonate, sodium hydrogen-carbonate, sodium hydrogen-carbonate, sodium hydrogen-carbonate, potassium phosphias, triathylamine and the like to give compound (31).

(0235) The compound [31] obtained in the same manner as in the above-mentioned Production Method is reacted in an abobol solvent such as methanol, ethanol and the like or a mixed solvent of an abobol solvent and a solvent such as DMF, accidentifies, THE, chiloroforn, dichibornethene, styly accation, methylene chindred, subvent and the like in the presence of a base such as potassium cerbonate, sodium carbonate, Ithium hydroxide, sodium hydrides, sodium hydrides, and the like at room temperature or with heating for deprotection, and reacted with compound [33] obtained in the same manner as in Step 1 of Production Method 1-1 in the same manner as in Step 1 of Production Method 5 to give compound [33].

Step 3

ō

(0238) The compound [33] obtained in the same manner as in the above-mentioned Production Method was sub-jected to cyclization in a solvent such as DMF, actionatile, THF, chloroform, dichloromethane, othyl acetale, methylane chiefde, bluene and the life in the presence of a copper catalyst such as copper() locido and the like or a paliation catalyst such as paliadium(ii) chloride and the like at room temperature or with heating to give compound (II-2-15).

Production Method 8

Ğ

[0237] Formation of tmidazo[1,2-a]pyridine ring

8

8 defined above. wherein Role and Rolf are each independently alkyl, such as methyl, ethyl and the like, and other symbols are as

8

[0238] The compound [34] obtained by the above-mentioned Production Method or a conventional method is subjected to smide condensation with compound [35] in the same manner as in Step 3 of Production Method 1-1 to give compound [38].

EP 1 162 196 A1

[0239] The compound (36] obtained by the above-mentioned Production Method is reacted with Grignard reagent [37] obtained by a conventional method to give compound [38].
[0240] Attemptively, an acid halide of compound [34] may be used instead of compound [36].

õ

[0241] The compound [38] obtained by the above-mentioned Production Method is subjected to halogenation by a

conventional method to give compound [39].

[0242] For example, when Hal is a bremine atom, compound [38] is reacted with bremine under cooling or at room temperature in a servent such as DMF, acatenticle, THF, chloroform, dichloromethane, ethyl acatelle, toluene and the

like to give compound [39].

[10243] Alternatively, a halogenating agent such as hypohelite (e.g., hypochlorite and the like), N-bromosuccinimide and the like may be used natioad of bromine for halogenation

Step 4

ĕ

8 [0244] The compound [39] obtained by the above-mantioned Production Method is subjected to cyclization with compound [40] obtained by a conventional or known method (I/A-A8-4885) in the presence of a base such as potassium actionate, acidium carbonate, acidium give compound [II-2-16].

[0245] The Production Methods shown in the above-mentioned Production Methods 2 to 4 can be used for the synthesis of compounds other than benzimidazole of the formulas (i) and (ii), such as compounds (ii-2-15) and (ii-2-15). (10245) The compounds of the formulas (i) and (ii), and honds thereof of the present invention are explicated in detail in the following by wey of Examples. It is needless to say that the present invention is not limited by those Examples.

Example 1

ક

Production of ethyl 2-(4-(3-bromophonoxy)phenyl)-1-cyclohexylbenzimidazolo-5-carboxylate

[0247]

Step 1: Production of eithyl 4-chioro-3-nitrobenzoste
4-Chioro-3-nitrobenzosia edid (300 g) was dissolved in eithyl alcohol (1500 m)) and concentrated sutfuric ecid
4-Chioro-3-nitrobenzosia edid (300 g) was dissolved in eithyl alcohol (1500 m)) and concentrated sutfuric ecid
(100 m)) was added with ice-conig. The instrume was refluxed under heating for 7 hr. The reaction mixture was
poured into ice-cold water and the precipitated crystals were collected by filtration to give the title compound (332)

5, yold 97%). HANNR (300MHZ, CDCL) : 8.50(1H, d, J=2.1Hz), 8.18(1H, dd, J=8.4, 2.1Hz), 7.83(1H, d, J=8.4Hz), 4.43(2H, q,

→7.5H2), 1.42(3H, 1, →7.5H2)

Step 2: Production of entyl 4-cyclohexylamino-3-nitrobenzoste

Step 2: Production of entyl 4-cyclohexylamino-3-nitrobenzoste

Step 2: Production of entyl 4-cyclohexylamino (30 g) obtained in the previous step was dissolved in scretonitrile (1500 ml),

Entyl 4-chtors-3-nitrobenzoste (300 g) obtained in the previous step was dissolved in scretonitrile over and cyclohexylamine (220 g) and triatilylamine (165 g) were added. The mixture was refluxed under heating overnight. The reaction mixture was poured into loc-cold water and the precipitated crystals were collected by tiltration to give the title compound (400 g, yield 84%).

H-NMR (300MHz, CDCt₃): 8.87(1H, d, Jac.1H2), 8.35-8.45(1H, m), 8.02(1H, dd, Jac.1H2), 6.87(1H, d. Jac.1H2).

8

J=9.1Hz), 4.35(2H, q, J=7.1Hz), 3.65-3.50(1H, m), 2.14-1.29(1DH, m), 1.38(3H, t, J=7.1Hz)

Step 3: Production of ethyl 3-emino-4-cyclohexylaminobenzosta

Step 4-cyclohexylamino-3-nitrobenzosta (400 g) obsinedad in the previous step was discoved in ethyl scenario (504 wet, 40 g) was added. The mixture was (1500 ml) and ethyl alcohol (500 ml), and 7.5% patiadium carbon (50% wet, 40 g) was added. The mixture was hydrogenated for 7 hr et atmospheric pressure. The catalyst was filtered off and the filtrate was concentrated under hydrogenated for 7 hr et atmospheric pressure. The catalyst was filtered off and the filtrate was concentrated under

8

reduced pressure. Dilsopropyl ether was added to the residue and the precipitated crystals were collected by filtration to give the title compound (289 g., yield 80%).

11-ANNR (300MHz, CDCl₃): 7.57(14, dd, J=8.4, 1.9Hz), 7.41(14, d, J=1.9Hz), 6.59(14, d, J=8.4Hz), 4.30(24, q, J=7.1Hz), 3.40-3.30(14, m), 2.18-2.02(24, m), 1.38-1.16(81, m), 1.35(314, 1, J=7.1Hz)

8tep 4: Production of ethyl 3-(4-(3-bremophenoxy)benzoy)lamino-4-cyclohexylaminobenzostie

layer was washed with saturated brine, dried over anhydrous magnesium sulfate, and concentrated under reduced pressure. Dictivit ether was added to the residue for crystalitzation and the crystals were collected by titration to dimothylomennide (catalytic amount) were added. The mixture was stirred for 4 hr at reom temperature. The reaction mixture was someontenine (160 ml). The re-sutting solution was concentrated ender reduced pressure and dissolved in dishiptormentane (160 ml). The re-sutting solution was added dropwise to a solution of ethyl 3-amino-4-cycloheayleminobanizatio (65 g) detained in the provious step in dichloromethane (500 mt) and triethylamine (71 mt), and the mixture was stirred for 1 hr at room temperature. The reaction mixture was pourod into water and extracted with dichloromethane. The organic 4-(3-Bromophenexy)benzoic eald (74 g) was dissolved in chloroform (500 ml), and exaly chlorido (33 ml) and

give the tibe compound (128 g. yield 95%).

HANIRI (390MHz, CDC4); 8.00-7.79(H4, m), 7.86(1H, pra), 7.37-7.16(3H, m), 7.13-6.86(3H, m), 6.72(1H, d. Jac.7Hz), 4.66(1H, bra), 4.26(1H, bra)

Step 8: Production of ethyl 24(4:3-bromophenoxy)phenyl)1-cyclohoxylbanzimidazole-6-carboxylate Ethyl 34(3-bromophenoxy)ebenzyl[artino-4-cyclohoxylatembenzoate (129 g) obtained tin the previous step Ethyl 34(3-bromophenoxy)ebenzoyl[artino-4-cyclohoxylatembenzoate (129 g) obtained tin the previous step was suspended to aceit aceit (600 ml) and tho resulting suspenden was offluxed under healting for 3 hr. The reaction mixture was concentrated under reduced pressure. Weter was added to the residue end the procepitated crystals were collected by filtration to give the title compound (128 g, 1914 89%).

1-4-MRR (300MHz, CDCQ); 8.5 (11H, d, J=8.7Hz), 1.60(1H, d), J=8.4, 1.5Hz), 7.57(1H, d, J=8.4Hz), 7.63(2H, d, J=8.7Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.53-7.21(3H, m), 7.17(2H, d, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 2.43-2.22(2H, J=8.7Hz), 7.14(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 4.42(2H, d, J=7.2Hz), 4.38(1H, m), 4.43(2H, d, J=7.2Hz), 4.38(1H, m)

5

Production of 2-(4-(3-bromophenoxy)phenyl)-1-cyclohoxylbenzimidezole-5-carboxytic acid

[0246] Eltryl 2-{4-(3-bromophonoxy)phenyl)-1-cyclohaxylbonzimidazzole-5-carboxylato (1.0 g) obtainod in Example 1 was affasolved in totrahydrofuran (10 ml) and athyl alcohol (10 ml), and 41 was dism hydroxide (10 ml) was added. The mixture was encountrated under reduced pressure and water was added to the residue. The mixture was cancentrated under reduced pressure and water was added to the residue. The mixture was additied with 6th hydroxibore add and the precipitated crystals were collected by fittration to give the title compound (0.9 g. yield 89%).

netting point: 255-256°C

8

'H-NMR (300Mhz, DMSO-d_e): (12.75(1H, 5x8), 8.24(1H, 8) , 7.88(1H, d, J=8.7Hz), 7.88(1H, d, J=8.7Hz), 7.71(2H, d, J=8.8Hz), 7.47-7.34(3H, m), 7.24(2H, d, J=8.8Hz), 7.20(1H, m), 4.31(1H, m) , 2.38-2.18(2H, m), 2.02-1.78(4H, m),

ô

Production of ethyl 1-cyclohexyl-2-(4-hydroxyphenyl)benzimidezole-5-cerboxylete

(0249) Ethyl 3-emino-4-cyclohexylaminobenzoate (130 g) obtained in Example 1, Step 3, and methyl 4-hydroxyben-zimidate hydrochloride (139 g) were added to methyl atcohol (1500 ml), and the mixture was refluxed under healing for 4 hr. The reaction mixture was allowed to cool end the precipitated crystats were collected by filtration to give tho

Ġ illie compound (131 g, yield 72%).
14-MMR (300MHz, CDCQ): 10.02(1H, bm), 8.21(1H, d, .=1.4Hz), 7.93(1H, d, .=8.8Hz), 7.83(1H, dd, .=8.8, 1.4Hz),
7.48(2H, d, .=8.8Hz), 8.53(2H, d, .=8.6Hz), 4.39.4.23(1H, m), 4.33(1H, d, .=7.0Hz), 2.35-2.18(2H, m), 1.88-1.79(4H, m),
1.70-1.80(1H, m), 1.48-1.18(3H, m), 1.35(3H, t, .=7.0Hz)

8

Production of ethyl 2-(4-(2-bromo-5-chlorobenzylexy)phenylj-1-cyclohexylbenzimidazole-5-carboxylate

ä In Exemple 3 were suspended in directly formands (200 m). Potasium carbonis (38 g) was added and the mixture was silved for 1 hr et 80°C with heating. The reaction mixture was silved to cool and then added to a mixed solven; of water-cityl secrets. The proceiptated crystals were active to y fination to give the title campound (50 g, yield 64%). of water-othyl eccisto. The prociplisted crystals were collected by filtration to give the title compound (50 g, yield 64%) "H-NMR (300MHz, CDCl_t): 8.50(1H, d, J=1.4Hz), 7.97(1H, dd, J=8.6, 1.4Hz), 7.70-7.57(5H, m), 7.20(1H, dd, J=8.4 (0250) 2-Bromo-5-chlorobenzyi bromide prepared from 2-bromo-5-chlorotoluene (50 g), N-bromosuccinimide and NN-azobisisobutyrantitile, and ethyl 1-cyclohoxyl-2-(4-hydroxyphonyl)benzimidazole-5-carboxylato (50 g) obisinod

EP 1 162 198 A1

m), 1.42(3H, t, J=7.1Hz) 2.6Hz), 7.14(2H, d, J=8.7Hz), 5.17(2H, e), 4.46-4.30(1H, m), 4.41(2H, q, J=7.1Hz), 2.40-2.20(2H, m), 2.02-1.21(8H,

Example 5

Production of ethyl 2-(4-2-(4-chlorophonyl)-5-chlorobenzyloxyl-phenyl)-1-cyclohoxytbonzimidazele-5-carboxylate

6 (2031) Ethyl 24(4-(2-bromo-5-chlorobenzyloxy)phonyl)-1-cyclohexyloenzimidazole-5-carboxylata (49 g) obtained in Example 4, 4-chlorophemylbornibe seld (18 g) and totralide (ritphemylborsphile)psilladium (10 g) were suspended in 1,2-dimethoxysthean (800 ml), 8 sutrated equeous sedium hydrogencentonate soution (300 ml) was added and the 1,2-dimethoxysthean (800 ml). Saturated equeous sedium hydrogencentonate soution, some mixer and saturated or the characteristic materials. The organic layer was washed successively with saturated equeous sodium hydrogencentonate soution, water and saturated brine, dried over anthydrous magnesium sulfata, and concentrated under reduced pressure. The residue was purified by silica gail flash chromologisphy (developing solvent, chloroform;shy) accesse = 97.3). Ethyl screeks and dileopropyl other were flash chromologisphy (developing solvent, chloroform;shy) accesse = 97.3). Ethyl screeks and dileopropyl other were added to the resulting oil for crystallization and the resulting crystals were collected by filtration to give the title compound

(44 g. yleid 65%). ¹H-NNR (300MHz, CDClg) : 8.49(1H, d., 1=1.4Hz), 7.97(1H, dd, 1=8.8, 1.8Hz), 7.70-7.80(2H, m), 7.56(2H, d, 1=8.7Hz), 4.95(2H, e), 4.48-4.28(1H, m) , 4.40(2H, m) , 2.02-1.20(8H, m) , 1.41(3H, t, 1=7.1Hz)

Example 6

8

Production of 2-{4-{2-{4-chlorophenyl}-5-chlorobenzyloxyjphenyl}-1-cyclohexybenzimidazele-5-cerborylic ecid

[0252] Ethyl 2-(4-(2-(4-chlorophenyl)-5-chtorobenzyloxylphonyl)-1-cyclohasylbonzimidazole-5-carboxylate (43 g) obtained in Exemple 5 was treated in the same menner as in Exempte 2 to give the title compound (33 g, yield 76%). metting point: 243-244°C

2

FAB-Ms: 571(MH+)

14-NMR (300MHz, DMSO-d₆): 8.32(1H, s), 8.28(1H, d, J=8.8Hz), 8.05(1H, d, J=8.8Hz), 7.76-7.72(3H, m), 7.58-7.48 (6H, m), 7.40(1H, d, J=8.3Hz), 7.24(2H, d, J=8.8Hz), 5.11(2H, s), 4.38(1H, m), 2.40-2.15(2H, m), 2.15-1.85(2H, m), 1.95-1.75(2H, m), 1.75-1.55(1H, m), 1.55-1.15(3H, m)

೪

č

Production of othyl 2-{4-{2-bromo-5-methoxybenzyloxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate

[0253] Ethyl 1-cyclohexyl-2-(4-hydroxyphenyl)benzimidezole-5-carboxylate obtained in Example 3 and 2-bromp-5-methoxybonzyl bromide were treated in the same manner as in Example 4 to give the title compound (59 g).

Example 8

â

Production of athyl 2 -{4 -{2-(4-chlorophenyl}-5-methoxybenzyloxy}-phenyl}-1-cyclohoxytbonzimidazole-5-carboxylate

Ġ [0254] Ethyl 24-4(2-bromo-5-methoxybenzyloxy)phenyl1-1-cyclohexybenzimidazoio-5-cerboxylate obtained in Example 7 was treated in the same manner as in Example 5 to give the till compound (48 g. yleid 77%). I+NMR (300MHz, COCI); 5-8-8(I)H, d. 1-4 Hz), 7-97(IH, d.d. 1-86, 1-4Hz), 7-8(IH, d. 1-86, Hz), 7-8(ZH, d. 1-87-Hz), 7-97(ZH, d. 1-86, Hz), m), 1.83-1.73(1H, m), 1.42(3H, 1, J=7.1Hz), 1.41-1.25(3H, m)

8

Production of 2-(4-(2-(4-chlorophonyl)-5-mothoxybonzyloxylphenyl)-1-cyclohoxylbonzimidazole-5-carboxylic acid

[0255] Ethyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)phenyl)-1-cyclohexylbenzimidazolo-5-carboxylata (52 g) obtained in Example 8 was treated in the same manner as in Example 2 to give the title compound (44 g, yield 89%). metting point: 248-249°C

8

H-NMR (300MHz, DMSO-d_e): 8.20(1H, 6) , 7.88(1H, d, 1=8.7Hz), 7.85(1H, d, 1=8.7Hz), 7.57(d, 2H, 1=8.6Hz), 7.48

(2H, d, 1=8.6Hz), 7.44(2H, d, 1=8.6Hz), 7.29(1H, d, 1=8.5Hz), 7.24(1H, d, 1=2.6Hz), 7.11(2H, d, 1=8.6Hz), 7.06(1H, dd, 1=8.5, 2.6Hz), 5.04(2H, e) , 4.26(1H, m), 3.83(3H, e) , 2.38-2.26(2H, m)

Example 10

Production of ethyl 1-cyclohexyl-2-(4-{(E)-2-phenylvinyl)phonyl)-benzimidazole-5-carboxylate

night at room temperature. The reaction mixture was ico-cooled and benzofuroxan (259 mg) dissolved in acetonitrile (2 mf) was added. The mixture was stirred for 7 hr at 60°C. The reaction mixture was ico-cooled. After 1N sodium hydroxide was eddad, ethyl accitie was added and the mixture was extracted. The organic layer was washed with water and saturated brine, dried over anhydrous magnesium sulfate, and concentrated under reduced pressure. The residue was purdled by slike gel flash chromstography (developing solvent, n-hexane:ethyl acetate = 4:1) to give the aloohol (6 ml) and trans-4-sülbenocsarbaldehyde (397 mg) was added under ice-cooling. The mixture was attrod over-Ethyl 3-amino-4-cyclohexylaminobenzoate (500 mg) obtained in Example 1, Step 3, was dissolved in methyl

ttle compound (540 mg, yield 63%). 1H-NNMR (300MHz, DMSO-4g): 8.28(1H, d, J=1.4Hz), 8.01(1H, d, J=8.7Hz), 7.90-7.80(3H, m), 7.75-7.65(4H, m), 7.50-7.25(6H, m), 4.35(2H, q, J=7.0Hz), 4.31(1H, m), 2.40-2.20(2H, m), 2.00-1.80(4H, m), 1.62(1H, m), 1.40-1.20(3H, m), 1. m), 1.36(3H, t, J=7.0Hz)

8 Example 11

Production of 1-cyclohexyl-2-(4-((E)-2-phenylvinyl)phenyl)-benzimidazole-5-carboxylic acid

i, [0257] Ethyl 1-cyclohaxyl-214-([E)-2-phenylvinyl]phenyl)-benzinidazole-8-carboxylate (127 mg) obtained in Example 10 was treated in the same manner as in Example 2 to give the title compound (116 mg, yield 97%). metting point: not lower than 300°C

¹H-NMR (300MHz, DMSO-d_g), 8.25(1H, s) , 7.98-7.28(13H, m) , 4.33(1H, brt), 2.41-2.23(2H, m), 2.03-1.78(4H, m), 1.71-1.58(1H, m), 1.49-1.20(3H, m)

g

Production of 2- (4-benzyloxyphenyl)-1-cyclopentylbenztmidazole-5-carboxylic acid

[0258] In the same manner as in Examples 1 and 2, the title compound (700 mg) was obtained

벊

J=9.0Hz), 5.16(2H, s), 5.03-4.89(1H, m), 2.41-1.63(8H, m) 14-MMR (300MHz, CDCl₃) : 8.60(1H, a), 8.04(1H, d, J=9.0Hz), 7.63(2H, d, J=8.4Hz), 7.51-7.32(6H, m), 7.14(2H, d,

Exemple 13

å

Production of 2-(4-benzyloxyphenyl)-1-cyclopentylbenzimidazole-5-carboxamide

- Ġ 8 (2259) 2-(4-Benzyloxyphenyl)-1-cyclopenylbenzimidazolo-5-carboxylic acid (700 mg) obtained in Exemple 12 was disablyed in dimethylismamido (10 ml), and simpolide (108 mg), 1-thityl-3-(3-dimethylismidopropyl)carbodimido hydrochierido (390 mg), 1-hydroxybenzoriazole (2075 mg) and triphylamine (0,3 ml) wore edded. The mixture was stirred overnight at room temperature. Water was added to the reaction mixture and the mixture was axirisated with eithyl acotate. The organic layer was washed successively with saturated aqueous sodium hydrogencarbonatio, water and saturated brine, driod over anhydrous medipassium sulfate, and concentrated under reduced pressure. Ethyl acotate and disappropyl other were added to the estitus for crystalization and the crystals were collected by litration to give the stiff a compound (571 mg, yield 81%).

FAB-Ms: 412(MH+) melting point: 232-233°C

1+NNMR (300MHz, CDCt₃); 8.29(1H, d, =1.5Hz), 7.88(1H, dd, J=8.5, 1.5Hz), 7.65-7.30(8H, m), 7.13(2H, d, J=8.8Hz), 5.16(2H, s), 4.93(1H, quint, J=8.8Hz), 2.40-1.80(8H, m)

8

g

EP 1 162 196 A1

Example 14

Production of 2-(4-bonzyloxyphonyl)-5-cyano-1-cyclopentylbenzimidazola

[0280] In the same manner as in Example 1, the title compound (400 mg) was obtained. FAB-Ms: 394(MH+) (E E 14-NMR (300MHz, CDCl₃): 8.11(1H, s), 7.68-7.30(8H, m), 7.13(2H, s) , 5.16(2H, s), 4.84(1H, quint, J=8.9Hz), 2.35-1.60

õ Example 15

Production of 2-(4-benzyloxyphenyl)-1-cyclopentylbenzimidazole-5-carboxamide oxime

š [0281] 2-(4-Benzyloxyphenyl)-5-cyeno-1-cyclopentythenzimidazole (400 mg) obtained in Example 14 was suspend-edin ethyl elcohol (3 m)) and water (1.5 m)), and hydroxylamine hydrochloidde (1.41 mg) and sodium hydrogenoarbonate (170 mg) were added. The mixture was refluxed under heating overnight. The reaction mixture was ellowed to cool and the precipitated crystals wore collected by filtration to give the title compound (312 mg, yield 71%).

14.NMR (300MHz, DMSO-dy): 8.20(1H, 9), 7.50-7.31(8H, m), 7.12(2H, d, ±8.7Hz), 5.15(2H, s), 4.94(1H, quint J=8.7Hz), 3.61(3H, e), 3.40(3H, e), 2.41-1.42(8H, m)

Example 18

10

Production of ethyl 1-cyclohexyl-2-(4-([4-(4-fluorophenyl)-2-methyl-6-thlazolyl)methoxy]phenyl)benzimidazole-5-carboxylate

[0262]

8

Step 1: Production of 4-(4-fluorophenyi)-5-hydroxymethyl-2-methylthiazola

Ethyl 4-(4-fluorophenyl)-2-methyl-5-thlazolocarboxylate (59 g) prepared by a known method (Chem. Pharm. Bull., 43(6), 947, 1985) was dissolved in tetrahydrofuran (700 ml). Lithium aluminum hydride (13 g) was added under ice-cooling and the mixture was stirred for 30 min. Neter (13 ml). 15% sectium hydroxide (13 ml) and water (39 ml) were added successively to the reaction mixture, and the precipitated insolutio materials were filtered off.

The filtrate was concentrated under reduced pressure to give the title compound (37 g, yield 71%). 14-NMR (300MHz, CDCly): 7.60(2H, dd, J=8.7, 6.6Hz), 7.11(2H, 1, J=8.7Hz), 4.80(2H, s), 2.70(3H, s)

Step 2: Production of Schloromethy4-4(4(loropheny)-2-methythlazole
4-(4-Fluoropheny)-5-hydroxymethy-2-methythlazolo (27 g) obtained in the provious step was dissolved in
chlorotom (500 ml), and thonyl chloride (24 ml) and pyridine (2 ml) were added. The mixture was stirred in 3 hr et room temperature. The reaction mixture was poured into to-cold water. The mixture was extracted with chic-roform, and washed with water and saturated brine. The organic layer was dried over sodium suifate, and con-

centrated under roduced pressure to give the title compound (28 g. yield 76%). 14.NMR (300MHz, CDCL): 7.67(2H, dd, J=8.8, 5.4Hz), 7.16(2H, t, J=8.7Hz), 4.78(2H, e) , 2.73(3H, e) Step 3: Production of ethyl Hoyolohexyk-2.(4-{(4-{d-fluorophenyi) -2-methyl-5-thiszolyilmethoxylphenyilbenzimidezole-5-carboxylate

6-Chloromethyl-4-(4-fluorophenyl)-2-mothylthlezole (28 g) obtained in the previous step and ethyl 1-cyclohexyl-2-(4-hydroxyphenyl)benzimidezole-5-carboxylate (38 g) obtained in Example 3 were treated in the same manner as in Example 4 to give the title compound (61 g, yield 100%). APCI-Ms: 570 (MH+)

14-NMR (300MHz, DMSO-d₃): 8.25(1H, d, J=1.5Hz), 7.97(1H, d, J=8.7Hz), 7.86(1H, dd, L=8.6, 1.6Hz), 7.7.4(2H, dd, J=8.6, 5.5Hz), 7.62(2H, d, J=8.7Hz), 7.33(2H, t, J=8.9Hz), 7.22(2H, t, J=8.9Hz), 5.41(2H, s), 4.34(2H, d, J=7.1Hz), 4.31(1H, m), 2.71(3H, s), 2.40-2.16(2H, m), 2.05-1.75(4H, m), 1.55-1.15(3H, m), 1.36(3H, t, J=7.1Hz)

Example 17

Production of 1-cyclohexyl-2-(4-((4-(4-(huorophenyl)-2-methyl-5-thiazolyl)methoxy)phenyl)benzimidazole-5-carboxylic

(1924) Ethyl 1-cyclohesyl-2144(1414-fluorophenyl)-2-methyl-5-thlazefylmethoxylphenylbenzimidazole-5-carboxy-tate (60 g) obtained in Example 16 was treated in the same manner as in Example 2 to give the title compound (39g, yield 65%).

netting point: 198-198°C

-RAP-Ma: Б42(MH+)
1-RAP-Ma: Б42(MH+)
1-H-MMR (300MHz, DMSO-d₃): 13.1(1H, bтв), 8.34(1H, s), 8.28(1H, d, J=8.3Hz), 8.08(1H, d, J=8.7Hz), 7.80-7.72(4H, m), 7.38-7.31(4H, m), 5.48(2H, s), 4.38(1H, m), 2.72(3H, s), 2.45-2.16(2H, m), 2.15-1.85(2H, m), 1.85-1.75(2H, m), 1.75-1.55(1H, m), 1.65-1.20(2H, m)

6

Production of ethyl 1-cyclohexyl-2-(2-fluoro-4-hydroxyphonyl)-benzimidazole-5-carboxylate

[0264] In the same manner as in Example 3, the title compound (50 g) was obtained.

8

Production of othyl 2-(4-[bis(3-fluorophonyl)methoxy]-2-fluorophonyl]-1-cyclohoxylbenzimidazole-5-carboxylate

6 [0265]

Step 1 : Production of 3,3'-diffuorobonzhydrol

The mixture was stirred for 10 min. The organic layer and water layer were separated. Water layer was extracted with ethyl acetals, and the combined organic layer was washed with 2H hydrochhote acid, saturated equeous sodium hydrogencarbonate and saturated brine. The organic layer was afrod over anhydrous magnesium sufface, filtered, and the selvent was evaporated off under reduced pressure to give the title compound (158.2 g. yeld 95%). 14-14MR (300MHz, CDCs): 7.31(2H, id, 1-7.9, 5.8Hz), 7.15.7.80(4H, m), 6.97-6.84(2H, m), 6.82(1H, d, 1-3.3Hz). To a stirred solution of magnesium strip (35.4 g) in THF (200 mt), todine strip was added and the mixture was heated with stirring under nitrogen stream until most of color of todine was disappeaned. A solution of 3-futoro-bromobenzene (230.0 g) in THF (1000 mt) was added dropwise over 2.5 fix while the temperature of the solution was maintained at 60°C. After the competition of the addition of the solution, the resulting mixture was refuxed for 1 hr with heating. The resulting Grignard solution was inco-cooled and a solution of othyl formatic (63.2 g) in THF 2.30(1H, d, J=3.3Hz) (200 ml) was added dropwise over 1 hr. After a stirring of the reaction solution for an additional 30 min, esturated aquoqua ammonium chioride solution (700 mt) was added dropwise with ice-cooling and water (300 mt) was added.

Step 2: Production of 3,3'-diffuorobenzhydryl chloride

8

ㅂ

To a solution of 3.3 difluorobenzhydrol (150.0 g) obtained in the previous step in toluene (400 ml), pyridine (539 mg) was added at room temperature. To the solution, thionyl chloride (89.1 g) was added dropwise over 1 hr at room temperature and the resulting solution was stirred for an additional 2 hr. The solution was heated so that was separated, and washed with water, saturated aqueous codium hydrogencarbonate and saturated brine. The organic layer was dried over anhydrous magnesium suffate, filtered, the solvent was ovaporated off under reduced pressure to give the title compound (158.2 g. yield 97%). the temperature of the solution was at 40° C, and then attreed for an additional 1.5 hr. Thionyl chiende (8.1 g) was added again and the mixture was stirred for 30 min. To the reaction mixture, water was added. The organic layer

Ĝ

H-NMR (300MHz, CDCh): 7.32(2H, td, J=8.0, 6.9Hz), 7.18-7.10(4H, m), 7.01(2H, tdd, J=8.2, 2.5, 1.2Hz), 8.05

Step 3: Production of ethyl 2-(4-[bis(3-fluorophenyt)methoxy]-2-fluorophenyt]-1-cyclohexylbenzimidazoto-5-cer-

Ethyl 1-cyclohoxyl-2-(2-fluoro-4-hydroxyphenyl)-benzimidszole-5-carboxylate (50 g) obtained in Example 18 and 3,3°-difluorobenzhydryl chloride (34 g) obtained in the provious step were treated in the same manner as in Example 4 to give the title compound (78 g, yield 89%).

5

8

H-NMR (300MHz, DMSO-dg): 8.24(1H, d, J=1.4Hz), 7.88(1H, d, J=8.7Hz), 7.88(1H, d, J=8.7Hz), 7.58(1H, t,

9

EP 1 182 198 A1

J=8.6H2), 7.50-7.40(6H, m), 8.82(1H, e), 4.34(2H, q, J=7.1H2), 3.85(1H, m), 2.20-2.10(2H, m), 1.80-1.80(4H, m) 1.6(1H, m), 1.35(3H, 1, J=7.2H2), 1.30-1.20(3H, m2)

Example 20

Production of 2-(4-(bis(3-fluorephenyf)methoxy)-2-fluorephenyf)-1-cyclohexylbenzimidazofe-5-carboxylic acid

[0288] Ethyl 244-(bts(3-fluorophenyl)melthory):2-fluorophenyl):1-cyclohery/benzimidazole-5-carboxylate (75 g) obtained in Example 19 was treated in the same mannor as in Example 2 to give the title compound (48 g, yield 62%). melting point: 242-242**C
FAB-Ma: 557(MH+)
FAB-Ma: 5

1.50-1.15(3H, 雨)

Example 21

Production of ethyl 1-cyclopentyl-2-(4-nitrophenyl)benzimidazole-5-carboxylate

[0267] In the same manner as in Example 1, the title compound (12 g) was obtained

Example 22

g

2 Production of ethyl 2-(4-aminophenyl)-1-cyclopentylbenzimidazote-5-carboxylate

(0268) Ethyl 1-cyclopantyl-2-(4-nitropheny)benzimidazole-5-carboxylate (12 g) obtained in Example 21 was dis-solved in tetrahydrolutan (200 m) and ethyl stoohle (50 m), 7.5% pafiadium canon (60% wor), 1 g) was added. The mbrure was hydroponated for 1 het atmosphorb pressure. The catalyst was litered off and the littrate was concen-trated under reduced pressure. Tetrahydrofuran was added to the residue to allow crystalization and the crystals were

collected by filtration to give the title compound (11 g, yield 88%).

14-NNR (300MHz, CDCL): 8.49(1H, d, J=1.3Hz), 7.95(1H, dd, J=8.5, 1.3Hz), 7.50-7.40(3H, m), 6.79(2H, d, J=4.6Hz),

4.97(1H, quint, J=8.9Hz), 4.40(2H, q, J=7.1Hz), 3.74(2H, brs), 2.40-1.60(8H, m), 1.41(3H, t, J=7.1Hz)

ä Example 23

g

Production of ethyl 2-(4-benzoylaminophonyl)-1-cyclopentylbenzimidazole-6-carboxylate

â tor 30 min at room temperature. The reaction mixture was concentrated under reduced pressure and water was added to the residue to allow crystalization. The crystals were collected by filtration to give the title compound (403 mg, yield dissolved in pyridine (3 mi) and chloroform (3 mi), and benzoyl chloride (127 mg) was edded. The mbrure was adred [0289] Ethyl 1-cyclopentyl-2-(4-aminophenyl)benzimidazole-5-carboxylate (300 mg) obtained in Example 22 was

14-NNR (300MHz, CDCb): 8.58(1H, s), 8.00(1H, d, J=9.0Hz), 7.84(2H, d, J=7.5Hz), 7.60-7.40(8H, m), 7.14(2H, d, J=7.5Hz), 4.84(1H, quint, J=8.7Hz), 4.41(2H, q, J=7.5Hz), 2.20-1.30(8H, m), 1.41(3H, t, J=7.5Hz)

Ġ

50

Production of 2-(4-benzeylaminophenyl)-1-cyclopentylbenzimidszele-5-carboxytic acid

(9270) Ethyl 2 (4 dearzydeminophony) 1-9yelopenytbanzimidazole-5-carboxylate (200 mg) abialmed in Example 23 was trosted in the same manner as in Example 21 to give the title compound (131 mg, yield 70%). methng point: not lower than 300°C FAD-Ms: 426(MH+) FAD-

Production of ethyl 2-(4-(3-chlorophenyl)phenoxy)phenyl)-1-cyclohexylbenzimidszele-5-cerboxylate

- [0271] Elhyl 2-[4-(3-bromophenoxy)phenyl}-1-cyclohexylbenzhnidazole-5-carboxylate (65 g) obtained in Example 1 and 3-chlorophenylboronic acid (23 g) were treated in the same manner as in Example 5 to give the title compound
- 5 (89 b, ylodd 85%).
 (89 b, ylodd 85%).
 (89 b, ylodd 85%).
 (14 +4NMR (300MHz, CDCty) : 8.51(1H, d, J=1.8Hz), 7.89(1H, dd, J=8.7, 1.8Hz), 7.71-7.55(4H, m), 7.51-7.43(2H, m),
 7.43-7.27(4H, m), 7.19(1H, d, J=8.4Hz), 7.12(1H, m), 4.41(2H, q, J=7.2Hz), 4.39(1H, m), 2.42-2.22(2H, m), 2.03-1.87
 (44, m), 1.78(1H, m), 1.42(3H, 1, J=7.2Hz), 1.39-1.29(3H, m)

Example 26

Production of 2-(4-(3-(3-chloropheny))phenoxy)pheny)-1-cyclohexy/benzimidazole-5-carbaxy/ic acid

- Ğ [0272] Ethyl 2-(4-(3-(3-chlorophenyl)phenoxy)phenyl)-1-cyclohaxylbenzimidazole-5-carboxylate (59 g) obtained in Example 25 was treated in the same manner as in Example 2 to give the title compound (43 g, yield 75%). melting point: 253-254°C
- g 14-NMR (300MHz, DMSO-d_d): 12.82(1H, bm), 8.24(1H, d, J=1.3Hz), 7.88(1H, d, J=8.7Hz), 7.88(1H, dd, J=8.7, 1.3Hz), 7.78(1H, e), 7.72(2H, d, J=9.7Hz), 7.70(1H, m), 7.64-7.42(5H, m), 7.25(2H, d, J=8.7Hz), 7.20(1H, m), 4.33(1H, m), 2.39-2.17(2H, m), 2.00-1.78(4H, m), 1.65(1H, m), 1.50-1.22(3H, m) FAB-Ms: 623(MH+)

ķ

Production of ethyl 2-[4-(3-acetoxyphonyloxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate

[0273] In the same manner as in Example 1, the title compound (87 g) was obtained

Production of ethyl 1-cyclohexyl-2-(4-(3-hydroxyphonyloxy)-phenyljbenzimidazole-6-carboxylate

- ä added. The mixture was stirred for 30 min at room temperature. The insoluble materials were filtered off and the filtrate was concentrated under reduced pressure. Water was added to the residue and the mixture was neutralized with 2N bydrochloric acid. The precipitated crystals were collected by filtration to give the title compound (78 g., yield 87%). 1-NNIMR (300MHz, DMSC-4g): 9.71(1H, e), 7.58(1H, d, Je8.7Hz), 7.87(1H, d, Je8.7Hz), 7.68 (2H, d, Je8.8Hz), 7.24 (1H, t, Je8.1Hz), 7.18(2H, d, Je8.6Hz), 6.63(1H, d), Je8.1Hz), 6.57(1H, d, Je8.1Hz), 6.51(1H, e), 4.38-4.23(1H, m), 4.36-4.23(1H, m), 1.36(3H, d), Je8.1Hz), 6.57(1H, d), Je8.1Hz), 6.57(1H, d), Je8.1Hz), 7.18(2H, d), 1.38-1.78(4H, m), 1.71-1.58(1H, m), 1.45-1.20(3H, m), 1.36(3H, m), 1.36(3H, m), 1.71-1.58(1H, m), 1.45-1.20(3H, m), 1.36(3H, m), 1.36(3H, m), 1.71-1.58(1H, m), 1.45-1.20(3H, m), 1.36(3H, m), 1.36(3H, m), 1.37-1.58(1H, m), 1.45-1.20(3H, m), 1.36(3H, m), 1.38-1.78(4H, m), 1.71-1.58(1H, m), 1.45-1.20(3H, m), 1.36(3H, m), 1.36(3H, m), 1.37-1.58(1H, m), 1.38-1.78(4H, m), 1.38-1.38(4H, m), 1.38-1 (0274) Ethyl 2-(4- (3-scetoxyphenyloxy)phenyl)-1-cycloherylbenzimidazole-5-carboxylate (87 g) obtained in Exam-ple 27 was dissolved in methyl sicohol (250 ml) and tetrahydrofuran (250 ml), and potassium carbonsis (31 g) was

- Ġ Production of ethyl 1-cyclohexyl-2-(4-[3-(4-pyridylmethoxy)-phenyloxy)phenyl)benzimidazole-5-carboxylate
- 8
- s), 4.47-4.31(1H, m), 4.42(2H, q, J=7.0Hz), 2.42-2.22(2H, m), 2.04-1.71(5H, m), 1.45-1.25(3H, m), 1.42(3H, 1, J=7.0Hz)

EP 1 162 196 A1

Production of 1-cyclohexyl-2-(4-[3-(4-pyridylmethoxy)phenyloxy]-phenyljbenzimidazeto-5-carboxylic acid

- [0278] Ethyl 1-cyclohasyl-2-(4-(3-(4-pyidylmethoxy)phonyloxy)-phenylibenzimidazole-5-carboxylate (80 g) obtained in Example 29 was treated in the same manner as in Example 2 to give the title compound (54 g, yield 75%). melting point: 235-237°C
- FAB-Ms: 520(MH+)
- õ 14-NMR (300MHz, DMSO-d₆): 8.58(2H, d, J=6.0Hz), 8.23(1H, s), 7.86 and 7.86(2H, ABq, J=8.7Hz), 7.66 and 7.17 (4H, A'8'q, J=8.7Hz), 7.44(2H, d, J=8.7Hz), 7.38(1H, t, J=8.0Hz), 8.80(1H, d, J=8.1Hz), 8.84(1H, s), 8.75(1H, d, J=8.1Hz), 6.22(2H, s), 4.40-4.22(1H, m), 2.40-2.19(2H, m), 2.00-1.80(4H, m)

Example 241

Production of methyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)-phenyl)-1-cyclohexylbenzimidezote

[0277]

- 8 Step 1: Production of 2-brame-5-mothoxybenzeidehyde
- 3-Methoxybenzaldehyde (16 g) was dissolved in acetic acid (75 m), and a solution of bromine (5.7 m) dissolved in acetic acid (16 m) was added dropwise. The mixture was stirred overnight at ream temperature and water (150 m) was added to the reaction mixture. The procipitated crystals were collected by filtration, washed with vater and dried under reduced pressure to give the title compound (21 g, yield 80%).

 14-ANMR (300MHz, CDC4): 10.31(14, s), 7.52(1H, d, 4-8.8Hz), 7.41(1H, d, 4-3.3Hz), 7.03(1H, dd, 3-8.8, 3.3Hz).

Step 2: Production of 2-(4-chlorophenyl)-5-methoxybenzaldehyde

- 2-Bromo-6-mothoxybenzaidehyde (10 g) obtained in the provious stop was treated in the same mothod as in Example 5 to give the title compound (11 g, yield 86%).

 14-NNRI (300MHz, CDCb): 9.82(14, e): 7.50(14, d, J-2.6Hz), 7.48-7.14(8H, m); 3.90(3H, e)

ä

25

- (30 ml). The solution was added dropwise to a suspension of sodium borehydrids (820 mg) in isopropyl alcohol (50 ml) and the mixture was stirred for 1 hr. The solvent was evaporated under reduced pressure and water was added to the residue. The precipitated crystals were collected by fittration and dried under reduced pressure. The Step 3: Production of 2-(4-chlorephenyi)-5-methoxybenzyi alcohol
 2-(4-Chlorephenyi)-5-methoxybenzaldehyde (10 g) obtained in the previous step was dissolved in tetrahydrofuran resulting crystals were recrystallized from a mixture of methanot and water to give the title compound (9.2 g, yield
- 14-NIMR (300MHz, CDCl₃): 7.37(2H, d, J=8.6Hz), 7.27(2H, d, J=8.6Hz), 7.17(1H, d, J=8.6Hz), 7.11(1H, d, J=2.6Hz), 6.88(1H, dd, J=5.6, 2.6Hz), 4.57(2H, e) , 3.86(3H, e) 8tap 4: Production of 2-(4-chlorophonyl)-5-mattoxybenzyl chlorido
- 2.(4-Chlonophenyl)-5-mathoxybenzyl alcohol (20 g) obtained in the previous step was disaolved in ethyl ace-tale (100 m) and pyridine (0.5 ml), and thorpyl chloride (11 ml) was added dropwise. The mixture was stirred for 1 hr. Water was added to the reaction mixture and the mixture was extracted with ethyl acctate. The organic layer was washed with water, atturated equeous sodium hydrogenicationate, water and saturated brine, dried over enhydrous magnesium suffate, and concentrated under reduced pressure. Isopropyl alcohol was added to the residue to allow crystellization. The resulting crystats worn collected by filtration and dried under reduced pressure to give the title compound (16 g, yield 74%).

à

- જ clohexytbenzimidazole-5-carboxylate 4.46(2H, s) , 3.86(3H, s) Step 5: Production of methyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)phonyl)-1-cy 14-NMR (300MHz, CDCl₃): 7.43-7.29 (4H, m) , 7.17(1H, d, J=8.6Hz), 7.05(1H, d, J=2.6Hz), 6.96-8.89(1H, m),
- 2-(4-Chlorophenyl)-6-methoxybenzyl chloride (4.0 g) obtained in the previous step and methyl 1-cyclohoxyl-2-(4-hydroxyphenyl)-5-enzimidazole-6-carboxylate (5.0 g) obtained in the same manner as in Example 3 were treat-
- 8 ed in the same manner as in Example 4 to give the title compound (6.0 g, yield 72%). 'H-NMR (300MHz, CDC4): 8.48(1H, s), 8.00-7.93(1H, m), 7.68-7.62(1H, m), 7.54(2H, d, J.=9.0Hz), 7.41-7.18(6H, m), 7.04-8.93(3H, m), 4.97(2H, s), 4.38(1H, m), 3.94(3H, s), 3.97(3H, s), 2.39-2.21(2H, m), 2.02-1.88(4H, m),

Example 242

Praduction of 2-(4-{2-(4-chiorophanyi)-5-mathoxybonzyloxyjbhonyi)-1-cyclohoxytbanzimidazolo-5-carboxylic acid

[0278] Mathyl 2-(4-(2-44-chlorophenyl)-5-methoxybenzyloxy)phenyl)-1-cyclohoxybenzimidazolo-5-carboxylate (5.0 g) obtained in Example 241 was treated in the same manner as in Example 2 to give the title compound (5.1 g, yield).

APCI-MB: 588(MH+)

5

14-NMR (300MHz, DMSO-4₆): 8.30(1H, d, J=1.4Hz), 8.24(1H, d, J=8.7Hz), 8.03 (1H, d, J=8.7Hz), 7.72(2H, d, J=8.7Hz), 7.51-7.39(4H, m), 7.34-7.18(4H, m), 7.11-7.03(1H, m), 5.08 (2H, s), 4.35(1H, m), 3.83(3H, m), 2.40-2.18 (2H, m), 2.10-1.98(2H, m), 1.93-1.78(2Hm), 1.72-1.18(4H, m)

Example 243

Ğ

Production of ethyl 2-(4-(3-(4-chtorophenyl)pyridin-2-yimethoxy)phenyl)-1-cyclohexylbenzimidazolo-5-carboxylate

[0278]

Step 1: Production of methyl 3-hydroxypicolinate

8

3-hydroxypicolinic acid (1.0 g)) was suspended in methenol (10, m) and concentrated sulfuric acid (10 m), was added. The mixture was refluxed under heating for 5 hr. The reaction mixture was toe-ocoled, neutralized with saturated acqueuts softlim hydrogencationate, and extracted with chlorolorm. The organic layor was washed with water and saturated brine, and dried over anhydrous magnestum sulfate. The solvent was ovaporated under reduced pressure to give the title compound (711 mg, yield 64%).

H-NMR (300MHz, CDC4): 10.63(1H, s), 6.28(1H, dd, J=3.7, 1.8Hz), 7.477.35(2H, m), 4.08(3H, s)

Step 2: Production of mithyl 3-(inthuoromothylsulfonyloxy)-pyridine 2-cantoxylsulfonyloxy)-pyridine Mothyl 3-hydroxypicoliant (70 mg) otherwise in provious step and triathylamine (0.77 mf) were disabved in dichloromothane (7 ml), and brilluoromothanosulfonic enhydrido (0.86 mf) was added under ice-cooling. The rescribe mixture was allowed to warm to room temperature and the mixture was allowed to warm to room temperature and the mixture was suffered to the reaction mixture was allowed to warm to room temperature and the mixture was selfered to the reaction mixture and the mixture was extracted with othyl acception. The organic layer was weenfold with saturated brine and dried over enhydrous magnesium sulfate. The solvent was evaporated under reduced pressure to give the title compound (1.2 g. yield 80%).

H-MARI (300MHz, CDC4): 8 906-27(1H, m), 7.75.7.70(1H, m), 7.83(1H, dd, J=8.2, 4.5Hz), 4.05(3H, s)

Step 3: Production of mathyl 3-(4-chlorophenyl)pyridine-2-carboxylate

8

3

Methyl 3-(influoromethylsulfonyloxy)pyridine-2-carboxylate (1.2 g) obtained in the previous step was treated in the same manner as in Example 5 to give the title compound (728 mg, yield 60%).

14-NIMR (200MHz, CDCly): 8.73-8.88(1H, m), 7.77-7.88 (1H, m), 7.49(1H, dd, J=7.8, 4.5Hz), 7.46-7.37(2H, m). 7.32-7.23(2H, m), 3.80(3H, e)

ŧ

ょ

hydroxide (1.8 ml) and water (4.8 ml). The insoluble materials were filtered off and the filtrate was concentrated under reduced pressure. The residue was purified by sities gel flash chromatography (developing selvent, n-hax-anorithyl acetate = 1:1) to give the title compound (208 may, yield 25%.

14-MMR (200MMz, CDC); 8.60(1M, dd, Ja-4.8, 1.5Mz), 7.60-7.65(1M, m), 7.40-7.48(2M, m), 7.29-7.38(1M, m), Step 4: Production of (3-(4-chioropheny/)pyridin-2-y/]methanol

Methyl 3-(4-chioropheny/)pyridin-2-carboxylete (720 mg) obtained in the previous step was dissolved in telrehydrofuran (10 ml) and the solution was los-cooled. Lithium sturninum hydride (180 mg) was added to the solution and the mixture was stirred for 1 hr. To the reaction mixture were edded successively water (1.8 ml), 15% sodium

7.27-7.20(3H, m), 4.63(2H, s)

Step 5: Production of ethyl 2-(4-(3-(4-chloropheny))pyridin-2-yimethoxy)phenyl}-1-cyclohexylbenzimidazola

8

Ġ

[3-(4-Chloropheny))pyridin-2-yi]methanol (200 mg) obtained in the previous atop was dissolved in chloroform (3 ml), and thlonyl chloride (0.13 ml) and pyridine (catalytic amount) were added. The mixture was stirred for 1 hr at room temperature and concentrated under reduced pressure. The residue was dissolved in dimathylio mamide manner as in Example 3 and potassium carbonate (250 mg) were added. The mixture was stirred for 3 hr with heating at 80°C. The reaction mixture was then allowed to cool. Water was added and the mixture was extracted with athyl acctate. The organic layer was washed with water and saturated brine, died over anhydrous magnesium (3 ml), and ethyl 1-cyclohexyF2-(4-hydroxyphenyl)benzimidazolo-6-carboxylete (232 mg) obtained in the same uliste and concentrated under reduced pressure. The residue was purified by silice get flash chromatography

9

8

EP 1 162 198 A1

(Govolophig advont, n-hoxanovalty) accisite = 1/2) to give the title compound (248 mg, yield 68%).
14-NNR (300MHz, CDCL) : 6.71(1H, dd, J=4.7, 1.4Hz), 8.48(1H, d, J=2.1Hz), 7.98(1H, d, J=10.2Hz), 7.71-7.92
(2H, m), 7.63(2H, d, J=6.7Hz), 7.45-7.34(5H, m), 7.04(2H, d, J=8.7Hz), 5.14(2H, e), 4.48-4.28(3H, m), 2.38-2.19
(2H, m), 7.63(2H, d, J=6.7Hz), 7.45-7.34(5H, m), 7.04(2H, d, J=8.7Hz), 5.14(2H, e), 4.48-4.28(3H, m), 2.38-2.19
(2H, m), 7.63(2H, d, J=6.7Hz), 7.45-7.34(5H, m), 7.04(2H, d, J=8.7Hz), 5.14(2H, e), 4.48-4.28(3H, m), 2.38-2.19

Example 244

Production of methyf 2-[4-(2-bromo-5-tert-butoxycarbonytbenzyfoxy)phenyl]-1-cyclohexytbenzimidazolo

[0280]

5

ö

Step 1: Production of tert-butyl 4-bromo-3-methylbonzosto
4-Bromo-3-methylbonzos eatd (25 g) was suspended in dichloromethene (200 mil), and oxalyl chiorida (12 mil) and dimohyldomatota eatd (25 g) was suspended in dichloromethene (200 mil), and dimohyldomanarida (catalyluc emount) were added. The mixture was dissolved in tetrahydrofluran (200 mil) and the solvent was evaporated under reduced pressure. The residue was dissolved in tetrahydrofluran (200 mil) and the solvent was evaporated under reduced pressure. The residue was dissolved in tetrahydrofluran (200 mil) and the solvent was so-cooled. To the solution was added dropwise a solution of potassium terr-butoxida dissolved and the solution was so-cooled. the mixture was extracted with eithyl accitate. The organic layer was washed with water and saturated brine, and dried over enhydrous magnesium surfate. The solvent was evaporated under reduced pressure to give the title compound (27 g. yeak 85%).

14. HAMAI (300MHz, CDCL): 7. 83(1H, d. J=2.2Hz), 7. 87-7. 53 (2H, m), 2.43(3H, s), 1. 58(9H, s)

15. Step 2: Production of mothyl 2-{4-(2-bramo-5-tent-buloxycarbonybonzyloxy)phonyli-1-cyclohoxy/benzhidazolain tetrahydrofuran (150 ml) and the mixture was stirred for 30 min. Water was added to the reaction mixture and

left-Buyl 4-bromo-3-methythenzeate (7.0 g) obtained in the provious step and methyl 1-cycloheayt-2-(4-hy-droxyphonyl)-bonzimidezolo-5-catboxylate (8.3 g) obtained in the same manner as in Example 3 were treated in the same manner as in Example 4 to give the title compound (8.8 g. yield 77%). H-NMR (300MHz, CDClg): 8.49(H, d. J-1.5Hz), 8.21(H, d. J-2.1Hz), 7.97(H, d. J-10.2Hz), 7.82(H, d. J-10.2Hz), 7.77-2.69(4H, m.), 7.18(CH, d. J-8.7Hz), 5.82(ZH, s), 4.38(1H, m), 3.95(JH, s), 2.40-2.23(ZH, m), 2.04-1.90(4H, m), 1.84-1.73(1H, m), 1.59(BH, s), 1.44-1.27(JH, m)

Example 245

g

ä 5-carboxylate Production of methyl 2-(4-(5-tent-butoxycerbonyl-2-(4-chlorophenyl)benzyloxylphenyl)-1-cyclohexylbenzimidezole-

[0281] Mothyi 2(4-(2-bromo-5-tent-butoxycarbony/benzybxy)phenyji-1-cyclohoxylbonzimidazolo-5-carboxylato (4.5 g) obtained in Example 244 was treated in the earne manner as in Example 5 to give the tible compound (3.6 g. yield 76%).

14-MMR (300MHz, CDC4₃): 8.48(1H, s), 8.27 (1H, d, J=1.8Hz), 8.04(1H, dd, J=7.9, 1.5Hz), 7.86(1H, dd, J=7.0, 1.5Hz), 7.65(1H, dd, J=7.86(1H, dd, J=7.86(1H, dd, J=7.86)), 4.89(2H, s), 4.43-4.28(1H, m), 7.05(1H, d, J=8.6Hz), 4.89(2H, s), 4.43-4.28(1H, m), 3.95(3H, s), 2.41·2.21(2H, m) , 2.02·1.89(4H, m) , 1.82·1.73(1H, m) , 1.62(8H, s) , 1.46·1.28(3H, m)

Example 246

8

Ĝ

Production of methyl 2-(4-(5-carboxy-2-(4-chlorophenyl)-benzyloxyjphonyl)-1-cyclohexybenzimidazolo-5-carboxylate mydrochloride

[0282] Methyl 2-(4-[5-tent-butoxycarbonyl-2-(4-chlorophenyl)-benzyloxylphenyl)-1-cyclohexylbenzimidazole-5-car-boxylate (3.5 g) obtained in Example 245 was disabled in dichloromethane (3.5 ml), and trilluoreactic acid (3.5 ml) was added. The mixture was altred for 1 hr at room temperature and the reaction mixture was concentrated under roduced pressure. The residue was disabled in othyl acotte, and 4N hydrochloric acid-othyl acotte was added. The prepipitate crystate were collected by filtration and dried under reduced pressure to give the title compound (3.3 g. ytold 97%).

8

2 14-NNR (300MHz, DMSO-d₃): 8.33(1H, d, J=1.5Hz), 8.29(1H, s) , 8.24(1H, d, J=1.8Hz), 8.09-8.00 (2H, m), 7.74(2H d, 1—8.8Hz), 7.81-7.44(5H, m) , 7.24(2H, d, 1—8.8Hz), 5.18(2H, s) , 4.38(1H, m), 3.83(3H, s) , 2.37-1,21(10H, m)

Production of methyl 2-(4-(2-(4-chlorophenyl)-5-methylcarbamoylbonzyloxy)phenyl)-1-cyclohoxylbonzimidazolo-

ĕ 5 sure. Water was added to the residue and the mixture was extracted with eithyl acetate. The organic layer was washed with water, saturated aqueous sodium hydrogencarbonate and saturated brine, and dried over anhydrous magnosium suffats. The solvent was evaporated under reduced pressure and the residue was crystalized from eithyl acetate and (0283) Mathyl 2-(4-(5-carboxy-2-(4-chlorophemy)benzyloxy)phemyl-1-speciohexy/denz/midazole-5-carboxy/sio hydrochtoride (400 mg) obtained in Example 264 was suspended in dishloromenthene (5 ml), and oxally chloride (0.08 ml) and dimethylicomentide (catalytic arount) were added. The mixture was stimed for 2 hr at room temperature. The reaction mbture was concentrated under reduced pressure and the residuo was dissolved in dishloromentane (5 ml). disopropyl ether. The crystals were collected by filtration and dried under reduced pressure to give the title compound rahydrofuran (5 ml) under lce-⇔ooling. The resction mixture was stirred for 1 hr and concentrated under reduced pres-The resulting solution was added dropwise to a mixed solution of 40% aqueous methylamine solution (5 ml) and tet-

(333 mg, yield 68%). 14-NNR (300MHz, COCk), : 8.47(1H, s), 8.08(1H, d, J=1.8Hz), 7.88(1H, dd, J=8.6, 1.5Hz), 7.82(1H, dd, J=8.2, 2.2Hz), 7.84(1H, d, J=8.8Hz), 7.54(2H, d, J=9.0Hz), 7.44-7.31(5H, m), 8.98(2H, d, J=9.0Hz), 8.35-6.28(1H, m), 5.00(2H, s), 4.36(1H, m), 3.95(3H, s), 3.05(3H, d, J=4.8Hz), 2.40-1.24(10H, m)

Example 248

2

Production of 2-(4-(2-(4-chiorophenyl)-5-methylcarbamoylbonzyloxy[phenyl]-1-cyclohexylbenzimidazolo-

t

8

[0284] Mothyl 2-(4-[2-(4-chbrophenyl)-6-methylcerbemoy/bonzyloxy]phenyl)-1-cyclohexylbenzimidezole-6-cerbox-ylata (150 mg) obtained in Example 247 and tetrahydrofuran (2 ml) were treated in the same mannor as in Example 2 to give the title compound (141 mg, yield 80%).

[0283] In the same manner as in Examples 1-30 and 241-248, and optionally using other conventional methods, where nocessary, the compounds of Examples 31-240, 249-327, 701 and 1001-1559 were obtained. The chemical structures and properties are shown in Table 1 to 177 and 165 to 212. 8.05-7.90(2H, m), 7.70(2H, d, J=8.6Hz), 7.56-7.43(5H, m), 7.21(2H, d, J=8.6Hz), 5.14 (2H, e), 4.34(1H, m), 2.81(3H 14-NMR (300MHz, DMSO-dg): 8.85-8.58(1H, m), 8.27(1H, d, J=1.5Hz), 8.21(1H, d, J=8.2Hz), 8.15(1H, d, J=1.5Hz) 1-4.5Hz), 2.39-1.19(10H, m)

t

Example 601

Production of methyl 2-[4-[2-(4-chlorophenyl)-5-methoxybenzyloxy]-phenyl]-1-cyclohexyl-1H-indole-5-carboxylete

[0288]

ŝ

t,

Step 1: Production of methyl 3-bromo-4-cyclohexylaminobenzoato

The reaction mixture was poured into 10% aqueous citic acid solution (100 mi) and extracted with shipty accrate (100 mi). The organic tayor was washed with water (50 mi) and saturated brine (50 mi), and dried over actium suifate. After (fitration, the solvent was evaporated under reduced pressure and the residue was purified by silica gail flash chromatography (developing solvent, n-hoxanezathyl scatate = 10:1) to give the title compound (2.6 g. yield 82%). 3-Bromo-4-fluorobenzolo acid (2.0 g) was dissolved in methanol (20 ml) and concentrated sulfunc soid (2 ml) was added. The mixture was refluxed for 3 hr. The reaction mixture was purped into los-cold water and extracted with ethyl accetate (60 ml). The organic beyer was washed with water (30 ml) and selurated brine (30 ml), and officid over sodium sulfate. After filtration, the solvent was evaporated under reduced pressure. The residue was dissolved over sodium sulfate. After filtration, the solvent was evaporated under reduced pressure. In dimethyl sulloxide (20 ml) and cyclohexylamine (10.3 ml) was added. The mixture was stirred overnight at 120°C

`н-NMR (300MHz, CDCl₃) : 8.10(1H, d, J=1.9Hz), 7.83(1H, dd, J=1.9Hz, 8.8Hz), 8.69(1H, d, J=8.7Hz), 4.73(1H,

8

8

brd, 4=7.3Hz), 3.85(3H, s), 3.38(1H, m), 2.10-2.00(2H, m), 1.90-1.20(8H, m)

Step 2: Production of 4-chloro-2-(4-lodophenoxymethy)-4-methoxybpheny,

4-lodophenic (6, 0) was disaphored in easione (50 m), and poteastim carbonate (4.7 g) and 4'-chloro-2-chloromethy-4-methoxybiphenyi (8.0 g) obtained in Example 241, Step 4 were added. The mixture was refluxed for

EP 1 162 196 A1

10 hr. The reaction mixture was concentrated and 4N equeous sodium hydroxide solution (50 mf) was added. The precipitated crystals were collected by filtration, washed with water, and dried under reduced pressure to give the title compound (10.0 g, yield 98%).

d, J=8.5Hz), 6.62(2H, d, J=8.9Hz), 4.84(2H, s), 3.85(3H, s) `H-NMR (300MHz, CDC\3) : 7.52(2H, d, J=8.9Hz), 7.35(2H, d, J=8.5Hz), 7.27-7.20(3H, m) , 7.12(1H, s) , 8.95(1H,

Step 3: Production of [4-(4'-chloro-4-mathoxybiphenyl-2-yimathoxy)phenylethynyljtrinethylsllane 4-Chloro-2-(4-iodophenoxymathy)-4-mathoxybiphenyl (7.0 g) obtained in the previous step was dissolved in aceionnitie (36 m), index dimentify)-4-mathoxybiphenyl (7.0 g) obtained in the previous step was dissolved in aceionnitie (36 m), index dissolved (2.8 g), indexid-(riphenyphosphina)pallation complax (1.8 g), copport() loadde (0.8 g) and intethylamine (50 m) were added. The mixture was attirad overnight at room temperorganic layer was washed with water (30 ml) and esturated brine (30 ml) and dried over sodium suffate. After (litration, the solvent was evaporated under reduced pressure and the residue was purified by silica gol flash ature and concentrated. Water (30 ml) was edded and the mbrure was extracted with ethyl acetate (50 ml). The

Step 4: Production of methyl 3-(4-(4-chloro-4-methoxybiphenyl-2-ylmethoxy)phenylethynyl]-4-cyclohexylemi-

5

g 2 g step was disastwoid in mathenoi (50 m) and othorotom (50 m)), and potessium carbonais (2.5 5) was added. The intritute was affired for 3 hr at room temperature and concentrated. Meatr (30 m)) was added and the mixture was extracted with othyr scenario (50 m). The organic layer was washed with water (30 m) and assurated brine (30 m), and dried over addition suitate. After fitterion, the solvent was organized under reduced pressure to give whits crystals (3.6 g). The white crystals (2.3 g) were dissolved in accionatific (10 m), and methyl 3-bronce-4-gobiner. ylaminoberscels (1.0 g) obtained in Step 1, terraksigtrophenylphosphhapjasiladium complex (0.4 g), expper(1) todido (0.1 g) and triothylismine (10 ml) were added. The mixture was stirred overnight at 100°C and concentrated under reduced pressure. Water (50 ml) was added and the mixture was extracted with athyl assetta (50 ml). That organic layer was washed with water (50 ml) and saturated brine (50 ml), and dried over sodium satista. After illitration, the solvent was evaporated under reduced pressure and the residue was purified by silica gal flash 8.85(2H, d, J=8.9Hz), 8.59(1H, d, J=8.8Hz), 6.07(1H, bm), 4.91(2H, s), 3.88(3H, s), 3.85(3H, s), 3.42(1H, m), 2.15-2.00(2H, m), 1.80-1.20(8H, m) chromatography (devoloping solvent, n-hexana-ethyl acetate = 8:1) to give the title compound (0,9 g, yiold 49%) 'H-NMR (300MHz, CDCi.j.: 8,03(1H, s), 7.84(1H, d, J=8.7Hz), 7.42-7.22(7H, m), 7.15(1H, s), 8.85(1H, d, J=8.2Hz) step 8: Production of methy/ 2-(4-Chlorophenyi)-5-methoxybenzyloxyjphenyi)-1 حyclohexyi-1H-indois-5-car [4-(4"-Chlore-4-methoxybipherryl-2-yimethoxy)pherrylethynyl]-trimsthytsilane (5.1 g) obtained in the previous

å Methyl 3. (4-(4'-chloro-4-methoxybiphenyl-2-yimethoxylphenylethynyl-4-cyclohexylaminobenzoete (0.5 g) obtained in the provious step was dissolved in Ni-dimethylformamide (5 m), and export(1) foldide (0.17 g) was added. The mixture was retinued to: 3 hr at 190°C. The hoxburble methods were removed by Ilization, Valuer (10 m) was added and the mixture was extracted with ethyl excelate (30 mi). The organic layer was washed with water (10 mi) and adjusted brine (10 mi), and dried over sedium suitae. After (firstant, the solvent was exponented under reduced pressure and the redictive was purified by silice gol flash chromatography (developing solvent, n-bestancethyl accitate = 81) to give the title compound (0.27 g, yield 53%).

14-NURF (300MHZ, CDCB): 3.4(1H, a), 7.65(1H, d, J-8.8Hz), 7.40-3.8Hz), 7.40-7.16(H, m), 7.00-8.94

(3H, m), 6.48(1H, s), 4.85(2H, m), 4.18(1H, m), 3.93(3H, s), 3.88(3H, s), 2.45-2.25(2H, m), 1.95-1.20(6H, m)

à

Production of 2-(4-[2-(4-chlorophenyi)-5-methoxybenzyloxy]phenyi)-1-cyclohexyi-IH-Indole-5-carboxytic acid

obtained in Example 501 was treated in the same manner as in Example 2 to give the title compound (0.18 g, yield 71%). [0287] Methyl 2-(4-(2-(4-chlorophenyl)-6-methoxybenzyloxy]phanyl]-1-cyclohoxyl-1H-Indole-6-carboxylate (0.27 g)

8

14-NMR (300MHz, DMSO-d₂): 12.43(1H, brs), 8.20(1H, s), 7.76(1H, d, b-9.3Hz), 7.72(1H, d, b-9.0Hz), 7.50-7.20(8H, m), 7.07-7.03(8H, m), 8.63(1H, s), 6.01(2H, s), 6.13(1H, m), 3.63(9H, m), 6.25-2.25(2H, m), 1.35-1.10(8H, m) (20.88) in the same meanner as in Examples 601 and 602, and optionally using other convocational methods where necessary, the compound of Example 503 was obtained. The chemical structure and properties are shown in Table 207.

Example 601

Production of ethyl 2-(4-benzyloxyphenyl)-3-cyclohexyllmidazo[1, 2-ajbyridine-7-carboxylate

Step 1: Production of 4-benzyloxy-N-methoxy-N-methylbenzamide

benzeirlazole (3.5 g) and triethylamine (3.5 ml) were added. The mixture was attred overnight at room temperature. Water was added to the reaction mixture and the mixture was extracted with ethyl acetale. The organic layer was washed auccessively with water, saturated aqueous sodium hydrogencarbonate, water and saturated brine, and compound (5.8 g. yield 84%). 'H-NMR (300MHz, CDCs): 7.22, 2H, d, J=8.8Hz), 7.28-7.46(5H, m) , 6.97(2H, d, J=8.8Hz), 5.10(2H, e) , 3.55(3H, dried over anhydrous magnesium sulfate.' The solvent was evaporated under reduced pressure to give the title 4-Benzyloxybenzoto edd (5.0 g) and N,O-dimethylhydroxylsmine hydrochloride (2.5 g) were suspended in dimethylfornemide (50 ml), and 1-(3-dimethylaminopropyl)-3-ethylcerbodiimide hydrochloride (5.0 g), 1-hydroxy-

Step 2: Production of 1-(4-benzyloxyphenyl)-2-cyclohexylethanone

Ì

ŭ

N-methylbonzamids (3, 8) obtained in the previous step was dissolved in ignary-driving (10 m) and the solution was added dropwide to the reaction mixture at room temperature. The mixture was stirred for 2 hr and saturated equations ammonitum citeridos solution was added to the reaction mixture. The mixture was stirred for 2 hr and saturated equations ammonitum citeridos solution was added to the reaction mixture. The mixture was extracted with diethyles shown the exponented under reduced pressure. The residue was purified by alike a got Itash chromatography (developing advent, n-hazanov:ethyla ecetate = 6.1) to give the title compound (3,8,9 yield 68%).

14.-HAMR (300MHz, CDCL): 7.93(2H, d, J=8.8Hz), 7.28.7.48(5H, m), 7.00(2H, d, J=8.8Hz), 5.13(2H, e), 2.78(2H, d, J=8.8Hz), 5.53(1H, m), 0.781.82(10H, m)

Step 3: Preduction of 1.4-denzy/soxyhenyly-2-dyromory-2-dyromory-2-dyromory was stirred for 10 mix at room temperature. Saturated aqueus ecitim hydrogencarbonate was added to the reaction mixture and the mixture was strated with dichlyl either. The organic layer was versided with water and assumented brine and off-sed over enhydrous magnatum surface, and the solvent was evaporated under roduced pressure. The readue was purified by alica get itsah chromatog-raphy (developing solvent, n-bezaraethyl acetate = 6:1) to give the title compound (888 ms), yield 65%).

14. HAMRI (300MHz, CDCL): 7.88(2H, d, J=8.8Hz), 7.28.7.48(6H, m), 7.02(2H, d, J=8.9Hz), 5.14(2H, s), 4.88(1H, J=8.8Hz), 1.98.3.2.1.1 (H), 1.98.3.2.1 (H Magnesium (470 mg) was suspended in tetrahydrofuran (2 mf) and cyclohexylmethyl bromide (3.4 g) was edded dropwise at room temperature. After the addition, the reaction mixture was stirred for 30 min at 80°C. The reaction mixture was allowed to cool and difuted with tetrahydrofuran (5 mf). Separately, 4-benzylexy-N-methoxy-

Ł

d, J=9.3Hz), 0.86-3.30(11H, m)

Step 4: Production of einyl 2 (4-berzyloxyphenyl)-3-cyclohexyllmidazo[1,2-a]pyridine-7-carboxylate
Ethyl 2-enrinopyridine-4-carboxylate (214 mg) prepared according to IP-A-8-48051, 1-(4-benzyloxyphenyl)2-bromo-2-cyclohexylatinanose (500 mg) oxioned in the previous atep and potasulare according to Servious atep and Service at Se pound (95 mg, yield 16%). APCI-MS: 455(MH+) Insoluble materials were littered off and the filtrate was concentrated under reduced pressure. The residue was purfiled by silica get flesh chromategraphy (developing selvent, n-hexane:eithyl acetate = 1:1) to give the title com-

14.NMR (300MHz, CDCL); 8.33(1H, s), 8.21(1H, d, 1=7.5Hz), 7.55(2H, d, 1=8.7Hz), 7.25-7.50(6H, m), 5.13(2H s), 4.41(2H, q, 1=7.1Hz), 3.25(1H, m), 1.41(3H, t, 1=7.1Hz), 1.16-2.00(10H, m)

Example 602

Production of 2-(4-benzyloxyphenyl)-3-cyclohexylimidazo (1,2-a)pyridine-7-carboxylic acid

[0290] Ethyl 244-benzyloxyphenyl-3-cyclohaxylimidazoj ("2-ejpyridine-7-carboxylate (95 mg) obtained in the previous step was treated in the same manner as in Example 2 to give the title compound (33 mg, 37%).

APCI-MS: 427(MH+)

14-NMR (300MHz, ĎMSO-d_s); 8.67(1H, d, 1∞7.3Hz), 8.08(1H, s), 7.25-7.58(8H, m), 7.13(2H, d, 1∞8.7Hz), 5.17(2H, s), 3.23(1H, m), 1.25-2.10(10H, m) [0231] The compounds shown in Tables 213 to 218 can be further obtained in the same manner as in Examples 1

EP 1 162 196 A1

to 701 or by other conventional method employed as necessary.

5 Purity Example No. >90% (NMR) 441 (0+1) 32 300MHs, CDC13 8, 51 (1H, d, J=1, 5Hz), 7, 98 (1H, d, J=8, 4Hz), 7, 61 (2H, d, J=8, 7Hz), 7, 56-7, 10 (6H, m), 7, 12 (2H, d, J=8, 7Hz), 5, 15 (2H, d, J=8, 7Hz), 5, 15 (2H, s), 4, 94 (1H, quint, J=9, 5Hz), 4, 41 (2H, q, J=7, 5Hz), 2, 44-1, 50 (8H, m), 1, 41 (3H, t, J=7, 5Hz) 18 NACR (8) ppm

	369 (#+1)	HS.
	>90% (NMR)	Purity
300Mhz, CDC13 7.81(2H, d., 196, 6Hz), 7.60 (2H, d., J=8, 8Hz), 7.51-7.21 (8H, m), 7.11(2H, d., J=8, 8Hz), 5.15(2H, s), 4.93(H, quin t., J=8, 8Hz), 2.36-2.30(2H, m), 2.09-2.04(3H, m), 1.75- 1.68(3H, m).		
IH NAR(6) ppm	No. 31	Example No.

	8	å	ô		t	8		t
Purity >90% (NMR)			Example No. 36	MS 427 (H+1)	Purity > 90% (NMR)			Example No. 35
	t, J=8. 8Hz), 2. 64(3H, s), 2. 40-1. 54(8H, m)	8. 40 (1H, d., J=1. 4Hs), 7. 95 (1H, dd, J=8. 6, 1. 4Hz), 7. 61 (2H, d., J=8. 7Hz), 7. 57-7. 30 (2H, d., J=8. 7Hz), 7. 57-7. 7Hz) 6H, db, 71. 13 (2H, d., J=8. 7Hz)	IH NUR(6) ppm				7. 91 (1H. s), 7. 59(2H, d, J=8 .7Hz), 7. 49-7. 30 (7H, n), 7. 11 (2H, d, J=8. 8Hz), 5. 15 (2H .s), 4. 19 (1H, quint, J=8. 8H	

8

8 Purity Example No. >90% (NMR) 456 (H+1) <u>پ</u> IH NAIR(\$) ppm 300MHz, CDC13 8. 20(1H, s), 7. 50-7.31(9H, s), 7. 12(2H, d, J=8.7Hz), 5. 16(2H, s), 4.94(1H, quint, J 78. 7Hz), 3.61(3H, s), 3.40(3H, s), 2.41-1.42(8H, m)

EP 1 162 198 A1

8

g

3 Puzity

>90% (NMR)

440 (N+1)

8

411 (2/+1)

3

8

Ġ

å

Example No.

జ

1H NMR(6) ppm

300MHz, CDC13
7, 84 (14, s), 7, 61 (2H, d, J=9
.0Hz), 7, 58 7, 30 (7H, m), 7, 12 (2H, d, J=9, 0Hz), 5, 16 (2H, s), 4, 94 (1H, quint, J=6, 7H, s), 3, 10 (6H, brs), 2, 40-1.5
0 (6H, m)

Example No.

မ္တ

IH NAR(8) ppm

300H±. CDC13 8.69(1H, s), 8.19(1H, d, J=9 .0H±), 7.62(2H, d, J=8, 7H±), 7.54(1H, d, J=9, 0H±), 7.48 -7.38(5H, m), 7.15(2H, d, J= 8.7H±), 5.17(2H, s), 4.98(1 H, quint, J=9, 0H±), 2.27-2, 07(6H, m), 1.82-1.78(2H, m)

ĸ Purity

483 (K+1)

Ē >90% (NMR)

37

1H NAR(6) ppm

Example No.

EP 1 162 196 A1

300Mis. CDC13 7. 66 (IH, s), 7. 61 (ZB, d, J=8 .8Hz), 7. 50-7. 28 (TB, m), 7. 12 (ZH, d, J=8. 8Hz), 6. 86 (IH .brs), 5. 15 (ZH, s), 4. 94 (IH .quint, J=8. 8Hz), 2, 97 (3H, s), 2. 29-1. 76 (8H, m).

IH NUR(8) ppm

Example No.

S

¥

Purity

426 (H+1)

>90% (NMR)

300M/b, CDC13 7, 72(14, 8), 7, 60-7, 35 (10H -0.1), 7, 10(2H, d.) J=8, 7/82), 5 -14 (2H, 8), 4, 90 (1H, quint, J=6, 8Hz), 2, 28-2, 19 (2H, m) -2, 19 (3H, s), 2, 19-1, 74 (6H

EP 1 162 198 A1

40

TH NAR(6) ppn

Example No.

>90% (NMR) 448 Q1+) 42 LH NAR(8) ppm

300MHz, DMS0
8. 11(1H, s), 7. 81 (1H, d, J=8
4Hz), 7. 72 (1H, d, J=8, 4Hz)
7. 65(2H, d, J=8, 4Hz), 7. 51
(2H, m), 7. 43(2H, m), 7. 37(1
H, m), 7. 29(2H, s), 7. 23(2H, s)
d, J=8, 4Hz), 5. 22(2H, s)
89(1H, quintet, J=9, 2Hz), 2
2-2. 0 (6H, m), 1. 7 (2H, m).

5

Purity

8

S Purity

>90% (NMR)

384 (1+1)

8

₹

Ġ

8

Example No.

39

1H NMR(8) ppm

300Miz, DMS0-46 7.84(IH, d, J=9, OHz), 7.79(2H, d, J=9, Thz), 7.52-7, 33(8H, m), 7.26(IH, d, J=9, OHz) 5.27(2H, 9), 4.92(IH, quin t, J=9, 3Hz), 2.19-1, 70(8H, m).

ŝ

ô

Example No.

Purity

>90% (NMR)

462 (H+1)

š

Puzity

>90% (NMR)

414(2+1)

S

MS 469 (M+1)	
Example No. 44	IH NMR(8) ppm
	300Mis. DESO-d6 12. 9(2H, bris), 8. 25(1H, s), 8. 00(2H, d, J=7. 8Hz), 7. 90(1H, d, J=8. 4Hz), 7. 74(1H, d, J=8. 7Hz), 7. 67(2H, d, J=9. Hz), 7. 62(2H, d, J=8. 1Hz), 7. 24(2H, d, J=8. 4Hz), 5. 32(2 H, s), 4. 88(1H, guint, J=9.0 Hz, 2. 25-1. 60(8H, m).
Purity >90% (NMR)	
MS 457(M+1)	

	MS 469 (M+1)
	Purity >90% (NMR)
300MHz, DMSO-d6 8. 33(1H, s), R, 08(1H, d, J=9 .0Hz), 7, 99(1H, d, J=9, 0Hz) 7. 49-7, 41(4H, m), 7, 33(2H d, J=8, 4Hz), 5, 22(2H, s), 4 .96(1H, quint, J=9, 0Hz), 2 25-1, 60(9H, m), 1, 30(9H, s)	
1H NMR(8) ppm	Example No. 43

Purity Example No. >90% (NMR) 453 (M+1) 481 (1+1) 47 300MHz, DMSO-d6 8. 33(1H, s), 8. 07(1H, d, J~8 8. 33(1H, s), 8. 07(1H, d, J~9, 0Hz) 7. 82-7. 72(6H, m), 7. 35(2H d, J~9. 0Hz), 6. 40(2H, s), 4 .95(1H, quint, J~8. THz), 2 .35-1. 60(8H, m). 1H NACR (&) ppm

(g)

3

10

20

S

Purity

>90% (NMR)

ŝ

Example No.

8

1H NMR(δ) ppm

EP 1 162 196 A1

Example No.

8

IH NMR(6) ppm

300MHz, DMSO-d6 B, 33(1H, s), 8, 07(1H, d, J=8 PMS), 7, 98(1H, d, J=8, Hz), 7, 198(2H, d, J=8, 4Hz), 7, 19(1H, d, J=3, 6Hz), 7, 19(1H, d, J=3, 1Hz), 2, 30-1, 60(8, Hz), 19(1Hz), 19(1Hz),

Purity

>90% (NMR) 447 (H+1)

ક્ષ

Purity

>90% (NMR)

443 (N+1)

z

Example No.

45

1H NAR(&) ppm

300Hfz, DMSO-d6
13. 4 (1H, brs), 8. 32 (1H, s), 6. 06 (1H, d, JeB, Thz), 7. 97 (1H, d, JeB, Thz), 7. 79 (2H, d)
14. 4. JeB, Thz), 7. 79 (2H, d)
158. 8Hz), 7. 56-7. 48 (4H, m)
1-8. 8Hz), 7. 56-7. 48 (4H, m)
1-8. 55 (1H, quint, JeB, shz), 5. 27 (2H, s), 4. 95 (1H, quint, JeB, shz), 2. 30-1. 60 (8H, m).

>90% (NMR)

414 (H+1)

IH NAR(6) ppm

EP 1 162 196 A1

Example No.

쥴 300MHz, DMSO-46 8. 93 (2H, d, J=6, 6Hz), 8. 35 (.H, s), 8. 06-8, 04 (3H, m), 7. 97 (1H, d, J=8, 7Hz), 7. 83 (2H, .d, J=6, 7Hz), 7. 38 (2H, d, J= 8. 7Hz), 5. 61 (2H, s), 4. 94 (1 H, quint, J=6, 7Hz), 2. 40-1. 60 (8H, m), 1-8.

300MHz, DMSO-d6 8. 33(1H, s), 8. 08(1H, d, J=8, 0Hz), 7. 18), 7. 99(1H, d, J=9, 0Hz), 7. 78(2H, d, J=8, 4Hz), 7. 39 (2H, d, J=8, 1Hz), 7. 32(2H, d, J=7, 7Hz), 7. 23(2H, d, J=7, 8Hz), 5. 22(2H, s), 4. 96(1H, quint, J=8, 0Hz), 2. 32(3H, s), 2. 32(3H, s), 2. 30-1, 60(8H, m).

ä

g

Purity 2

>90% (NMR)

470 (04+1)

25

Example No.

ä

8

ö

Purity 돐

>90% (NMR)

323 (X+1)

300Ms, puso-46 9.18(1H, t, J=5, 6Hs), 8.34(1H, s), 8.04(1H, d, J=9, 6Hs), 7.98(1H, d, J=6, 7Hs), 7, 62-7, 32 (7H, a), 5.27(2H, s), 4.95(1 H, quint, J=9, 0Hs), 3.99(2H d, a), 5.74s), 2.40-1, 60(8H

1H NAR(8) ppm

300Mha, DMSO-d6 8, 32(H, 4), 8, 05 (H, 4), J=8 8, 37(H, 8), 8, 05 (H, 4), J=8, 7(h), 7, 80(2H, 4, J=8, 4Hz), 7, 67 (1H, t, J=4, 5Hz), 7, 65 (H, t, J=4, 5Hz), 7, 45-7, 42(2H, th, J=4, 5Hz), 7, 45-7, 44(2H, th, J, 7, 35 (2H, d, J=8, 4Hz), 5, 3 1(ZH, s), 4, 96 (H, quint, J=9, 0Hz), 2, 30-1, 50 (8H, th,

G

Purity 5

>90% (NMR)

432 (M+1)

ដ

Example No.

Ġ

ŝ

ĸ

Purity

>90% (NMR)

427 (H+1)

K

g

80

50

IH NMR (&) ppm

8

300MHz, DMSO-d6 8. 31(1H, s), 8. 03(1H, d, J=9) .0Hz), 7. 93(1H, d, J=9, 0Hz) .7. 77(2H, d, J=8, 4Hz), 7. 31 (ZH, d, J=8, 7Hz), 5. 07(2H, s) .4. 94(1H, quint, J=8, 7Hz) .2. 45(3H, s), 2. 56(3H, s), 2 .2. 45(3H, s), 2. 56(3H, s), 2

51

1H NAR(6) ppm

8

8

3 Purity

>90% (NMR)

447 (2+1)

ŝ

å

Example No.

54

IH NAR (&) ppm

55

300MHz, DMSO-d6 12. 7(1H, brs), 10.0 (1H, s), 8. 22(1H, s), 7. 87(1H, d, J=8 .6Hz), 7. 69(1H, d, J=8.6Hz), 7. 53(2H, d, J=8.6Hz), 6. 96 (2H, d, J=8.6Hz), 4. 89(1H, q uint, J=9.0Hz), 2. 30-1. 60(8H, m). IH MIR(8) ppm

EP 1 182 198 A1

ဌ္ဌ

Example No.

8 6 8 8 8 6 6

MS 413 (H+)	
Example No.	57 1H NAR(6) ppm
	300MHz, DMSO-d6 10. 99 (31H, s.), 8. 26 (1H, s.), 8 .01-7, 86 (4H, m.), 7. 69-7, 59 (5H, m.), 7. 38 (2H, d.), 7-8. 7Hz), 4. 86 (1H, quínt, J-8. 7Hz), 2. 12-1, 90 (6H, m.), 1. 72-1. 69 (2H, m.)
Purity >90% (NMR)	
MS 462 (M+1)	

		5
	Purity Lang (NMB)	υŢ
300MAs, DMSO-d6 12.78(1H, br a), 8.24(1H, s), 7.88and7.7 2(2H, ABq, J=8, 6Hz), 7.66an d7.22(4H, AB q, J=8, 6Hz), 7.68(1H, s), 7.48-7, 42(3H, m), 5.24(1H, s), 4.88(1H, q) 4.11, J=8.8Hz), 2.30-1, 91(6H, m), 1.78-1.60(2H, m)		3
IH NAR(8) ppm	Example No. 55	Ε×
	والمرازون المرازون المرازية والمرازية والمرازي	I

Table 9

EP 1 162 198 A1

5	Purity		Example No.	
460 (H+1)	>90% (NMR)		No.	
¥+1)	(NMR)		59	
•		300H#, DMSO-66 10. 82 (1H, s), 8. 34 (1H, s), 8 14and7, 84 (4H, ABq, J=6, 4H s), 8. 06and7. 66 (4H, A'B' q, J=8. 6Hz), 8. 06-7. 88 (4H, m) 5. 5. 01 (1H, quint, T=6. 3Hz), 2. 35-2. 15 (4H, m), 2. 11-1. 9 6 (2H, m), 1. 80-1. 62 (2H, m)	IH.NMR(6) ppm	
				•

	494 (H+1)	SW
	>90% (NMR)	Purity
8.26-7, 72(H, m), 4.92(1H, 8.26-7, 72(H, m), 4.92(1H, quint, J=9,0Hz), 2.34-1,70 (6H, m), 1.75-1,61(2H, m)		- -
DISO-46	No. 58	Example No.

EP 1 162 198 A1

Table 10

EP 1 162 196 A1

돐 Purity Example ö. >90% (NMR) 532 (H+1) 62 300MBs, DMSO-d6 10. 6(1H, s), 8. 34(1H, s), 8. 10. 45(1H, s), 8. 34(1H, s), 8. 09-7. 98(4H, m), 7. 82(2H, d, J=8.7 Hs), 7. 50-7. 35(5H, m), 7. 20 -7. 17(2H, d, J=9. 0Hs), 6. 24 (2H, s), 5. 01(1H, quint, J=9. 3Hz), 2. 40-1. 60(8H, m). 1H NAR(8) ppm

Table 11

8 ů t 8 2 ô 8

> Purity Example No. >90% (NMR) 448 (J+1) 65 300Htz, DMS0-46 12. 64(IH, brs), 47, 2Hz), 7, 59 7, 80(IH, d, J=7, 2Hz), 7, 59 (IH, d, J=8, 7Hz), 7, 48-7, 30 (6H, m), 6, 11(2H, s), 5, 03(I H, quint, J=8, 7Hz), 4, 20-4, 05(2H, m), 3, 46-3, 90(3H, m) 2, 15-1, 60(12H, m) IH NAR(6) ppm

Purity ₹ Example ĕ. >90% (NMR) 427 QF+1) 2 300MHz, DASO-d6 12. 25 (1H, brs), 7, 70-7. 30 (9H, m), 7, 20 (2H, d, J=8, 7Hz) 7, 14 (1H, d, J=8, 4Hz), 5, 20 (2H, s), 4, 84 (1H, quint, J=6 .0Hz), 3, 66 (2H, s), 2, 30-1. 51 (8H, m) IH NAR(8) ppm

G

5

EP 1 162 196 A1

78

8

5

Purity

参90% (NMR)

457 (11+1)

ä

幺 Purity

>90% (NMR)

508 (H+1)

#

8

Ġ

å

Example No.

ය

IH NAR(&) ppm

Example No.

66

1H Núck (8) ppm

300MHz, MISO-46 10.59(iH, s), 8, 31(iH, s), 8 10.62H, d, J=8, 6Hs), 8, 03(i, H, d, J=8, 7Hz), 8, 00-7, 85(3, H, a), 7, 80(2H, d, J=8, 6Hs), 7, 41(2H, d, J=8, 2Hs), 4, 98(i, 1H, quint, J=8, 8Hs), 2, 71-1 1, 10(19H, m)

ক

300MHz, DMSD-d6 8. 27(H, s), 8. 14(IH, d, J=8 8. 27(Hz), 7. 96(IH, d, J=8. 4Hz) 7. 71(2H, d, J=9. 0Hz), 7. 51 (2H, d, J=6. 9Hz), 7. 48-7. 37 (3H, m), 7. 30(2H, d, J=98. 4Hz), 5. 25(3H, s), 4. 39(1H, m), 3. 27(3H, s), 2. 60-1, 95(6H, m), 1. 26-1.05(
2H, m), 5.

Ł

Purity

>90% (NMR)

443年1)

5

ä

5

Example No.

క్ష

1H NMR(6) ppm

300MHz, DMSO-d6 8. 32(1H, a), 8. 26(1H, d, J=8 .7m2), 8. 04(1H, d, J=8, 7Hz), .7.77(2H, d, J=8, 4Hz), 7. 52 (2H, d, J=6, 9Hz), 7. 46-7. 39 (6H, m), 5. 28(2H, s), 4. 38(1 H, m), 3. 71(1H, m), 2. 60-2. 1 5(2H, m), 2. 04-1. 96(4H, m), 1. 30-1. 20(2H, m).

å

		•	
SW	Purity	<u>*</u>	Example No.
441 (M+1)	>90% (NMR)		No.
+1)	NMR)		69
	Ę	300MHz, Du 9, 88 (1H, 6) 32 (1H, 6) 32 (1H, 6) 48q, J=9, 6 (4H, A' B' 2H, d, J=7, 2H, d, J=7, J=7, 8H2), 1, 96 (2H, n	1H NAR(8) ppm
		(SO-d6 (), 9. 42 (1H 8. 09and8. H2), 7. 81a H2), 7. 81a J-9. 2Hz) 8Hz), 7. 31 7.00 (1H, t (1H, quint 2. 17 (4H, m) 1. 183-1.	s) ppm
), s), 8. 02(2H, 02(2H, 7.50) (2H, t, J=7.8 J=8.7H 54(2H,	

MS 481 (M+1)	Purity >90% (NMR)	19-2	Example No. 68
		300MF, DMSO-d6 8.31 (1H d, J=1, 4Hz), 8.05 (1H, d, J=8, 6Hz), 7.96 (1H, d, J=8, 6Hz), 8.86-8.61 (4H, m) 7.51 (1H, d, J=6, 3Hz), 7.33 (2H, d, J=8, 8Hz), 5.28 (2H, s)), 4.94 (1H, quint, J=8, 8Hz)), 2.31-1.60 (8H, m)	1H NAR(8) ppm

	>90% (NMR)	Purity
300MHz, DMSO-d8 12.81 (1H, thes), 8.42 (1H, s) .7.90 (1H, d, J=8.5Hz), 7.80 .7.50 (6H, m), 7.44 (2H, d, l= 8.6Hz), 5.25 (2H, s), 4.88 (1 H, quint, J=8.8Hz), 2.30-1. 52 (8H, m)		. Š=°
IH NAR(8) ppm	67	Example No.

Table 13

EP 1 162 198 A1

Purity Example No. >90% (NMR) 483 (H+1) 72 1H NAR(8) ppm
300kHs, pisco-d6
8, 30 (1H, s), 8, 25 (1H, d, J=8
7, 76 (2H, d, J=8, 7hs), 7, 51
(2H, d, J=7, 2Hs), 7, 46-7, 33
(5H, m), 6, 27 (2H, s), 4, 36 (1, s), 2, 60-2, 25 (2H, m), 2, 1
6-2, 00 (2H, m), 1, 95-1, 85 (2H, m), 1, 20-1, 1
0 (2H, m), 0, 87 (9H, s).

MS 427 (1+1)	Purity . > 90% (NMR)		Example No.
			71
		300MHz, DMSO-d6 8. 31(11, 9), 8. 05 (11, d, J=8 .712), 7. 97 (111, d, J=8, 714x) 7. 76 (211, d, J=8, 614x), 7. 44 -7. 19 (711, m), 4. 94 (11, quin t, J=8, 81x), 4. 35 (211, t, J=6, 714x) .715), 3. 10 (211, t, J=6, 714x) ,2. 32-1, 60 (81, m)	1H NAR(8) ppm

g

ES.	Purity		Example No.
489 (И+1)	>90% (NMR)		No. 70
		300MB, DMSO-d6 8. 27 (1H, d, J=1. 2Hz), 8. 04 (.1H, d, J=8. 7Hz), 7. 94 (1H, d, J=8. 7Hz), 7. 72 (2H, d, J=8. 7 Hz), 7. 60-7. 20 (12H, m) 6. 74 (1H, s), 4. 92 (1H, quint, J=8 .9Hz), 2. 30-1. 58 (6H, m)	IH NACR (&) ppm

8

Table 14

EP 1 162 196 A1

EP 1 162 196 A1

Table 15

Example No.

2

LH NMR(8) ppm

8

ä

ß

8 ĸ 8 2 Purity Example S Purity Example No. No. >90% (NMR) >90% (NMR) 425 (M+1) 78 77 300MHz, DMSO-d6 8. 14(1H, s), 7. 79(1H, d, J=9 .0Hz), 7. 67(1H, d, J=8, 7Hz) 7. 40-7. 20(5H, m), 4. 89(1H .quint, J=8, 7Hz), 3. 54(2H, s), 3. 19-2. 90(3H, m), 2. 23-1. 69(14H, m) 300Miz, paso-d6 8, 21(1H, s), 7, 87 (1H, s), 7, 56and7, 43(4H, ABq, J=8, 1Hs), 7, 34-7, 16(5H, m), 4, 25(1 h, br., J=12, 5Hz), 3, 06-2, 9 2(4H, m), 2, 41-2, 17(2H, m), 1, 72-1, 5 8(1H, m), 1, 48-1, 15(3H, m) IH NMR(8) ppm 1H NACR (6) ppm

EP 1 162 196 A1

8

404 (24+1)

Purity MS

>90% (NMR)

476 (M+1)

9

Example No.

75

1H NMR(8) ppm

300MHz, DMSO-d6 12.80(1H, s), 8.26(1H, s), 7 .90(1H, d, 175, 2Hs), 7.76-7 .60(8H, m), 7.35(2H, d, 178, .412), 4.64(1H, quint, 178, 8 .412), 3.23(3H, s), 2.32-1, 90 (6H, m), 1.78-1.61(2H, m) 5

412(24+1)

8 4 6 8 8 6 6

SH	Purity		Example No.	
495 (H+1)	>90% (NMR)		٥.	
_	MR)		81	
		300MHs, 1MSO-d6 12.76 (H, brs), 8. 21 12.14 (H, rs), 8. 21 13. 14 (H, d), 149 (H, d Hs), 7. 85 (H, dd, J-e Hs), 7. 70-7. 55 (H, d), 1-8 (2H, d, J-es, 7Hs), 6. 2 14. 36-4, 15 (1H, m), 18 (2H, m), 2. 00-1, 7 18 (2H, m), 2. 00-1, 7 15 (3H, m)	1H NMR(8) ppm	
		1 (1H, d, J=8.6 1, J=8.6 1, 1, 4 1), 7, 23 15 (2H, 2 2, 39–2 1, 48–1		

Example No. 80 IH NRR(8) ppm 30MHz, DKS-7, 62 (1H, d, J=8 8.17 (1H, D), 7. 84 (1H, d, J=8 8.17 (1H, d), 7. 84 (1H, d, J=8, 1Hz), 5. 06-4, 81 (2H, d), 7. 48 (2H, d, J=8, 1Hz), 5. 06-4, 70 (2H, d), 7. 3.30-3, 12 (1H, D), 2.48-2. 31 (5H, D), 2. 15-1.60 (12H, D) Purity > 9.0% (NMR) BY AND IN THE CONTROL OF
--

	418 (M+1)	SH
	Purity > 9 0% (NMR)	Pu;
300Miz, DMSO-48 8. 15 (1H, e), 7. 81 (1H, d, J=8 4Hz), 7. 59 (1H, d, J=9, 0Hz) 7. 50-7 38 (5H, m), 5. 05 (1H , quint, J=9. 0Hz), 3. 85-2. 9 5 (3H, m), 2. 20-1. 65 (14H, m)		
IH NMR(8) ppm	Example No. 79	Ex
		١

EP 1 162 196 A1

Table 17

EP 1 162 198 A1

MS	Purity		Example No
441 (K+1)	>90% (NMR)		No.
		~	84
-		Insture of 1 somers (cis: trans=3:1) 300MHz, DMSO-d6 300MHz, DMSO-d5 300MHz, DMSO-d6 300, 1, 1, 1, 2, 2, 3, 4, 1, 7, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,	TH NOR(6) ppm

SW	ยูน	·	š .	E.
	Purity	_		Example No.
539 (X+1)	>90% (NMR)	\ \ \	5	0. 83
		. 5. 22 (2H, S), 4. 36 (1H, m), 2 . 50-1. 40 (10H, m), 1. 31 (18H , s).	13. 2(1H, brs), B. 30(1H, s), 8. 23(1H, d, J=8. 8Hz), B. 02(1H, d, J=8. 7Hz), 7. 74(2H, d, J=8. 6Hz), 7. 40-7. 33(5H, m)	1H NAR(6) ppn 300aHz, DASO-d6

MS · 603 (M+1)	Purity >90% (NMR)		
		300kHz, D&SO-d8 8. 27(1H, s), 8. 22(1H, d, J=8, THz) 7. 72(1H, s), 8. 22(1H, d, J=8, THz) 7. 69(2H, d, J=8, THz), 7. 60 7. 50(4H, m), 7. 45-7. 25(8H, m), 6. 75(1K, s), 4. 21-4, 23 (1H, m), 2. 39-2. 18(2H, m), 2. 10-1. 18 (4H, m)	

Table 18

3.048.18, 1.85.7-06 12.76 (1H, brs), 8. 16 (1H 12.76 (1H, brs), 8. 16 (4H, ABq.) 7.918.04, 7.442.046, 86 (4H, ABq.) 1.948, 6Hz), 7.39-7.29 1.94, 82 (2H, a), 4.35 (1 1.912, 2Hz), 2.35-2 1.91, 197-1, 76 (4H, m) 1.97-1, 56 (1H, m), 1, 45-1, 1 1.91	>80% (NMR)	Pur1ty
NAR(8)	NO 87	Example No.

MS 477 (H+1)	Purity >90% (NMR)		Example No. 86	
		300MHz, DMSO-66 12. 76 (H, d, J=7. 6Hz), 8. 22 (1H, s), 8. 12.7 (1H, d, J=7. 6Hz), 8. 02-7. 53 (10H, m), 7. 32 (2H, d, J=8. 7Mz), 5. 68 (2H, s), 4. 32 (1H, br.t, J=12. 2Hz), 2. 41-7. 20 (2H, m), 2. 01-1. 78 (4H, m), 1. 71-1. 56 (1H, m), 1. 50-1. 16 (3H, m)	1H NAR(8) ppm	

MS 491 (M+1)	Purity >90% (NMR)		Example No. 85
•	•	300M/s, DHSO-d8 3.25 (1H, s) , 8. 14-7, 83 (6H, s), 7. 77-7, 44 (5H, s), 7. 21 (s), 4. 44 (2H, s), 7. 31 (s), 4. 44 (2H, s), 4. 44 (2H, s), 4. 44 (2H, s), 4. 45 (2H, s), 4. 31 (1H, s), 5. 66 (2H, s), 5. 20 (2H, s), 5. 20 (2H, s), 5. 20 (2H, s), 7. 70-1, 56 (1H, s), 1. 45-1, 14 (3H, s)	IH NAR(&) ppm

Table 19

EP 1 162 196 A1

Purity

Example No. >90% (NMR) 531(H+1) 8 1H NMR(&) ppm 300MHz, MMSO-d6 8. 40-8. 20 (2H, m), 8. 04 (1H, d, J-8. 4Hz), 7. 65 (2H, d., J-8, d, J-9. 4Hz), 7. 50-7. 10 (12H, m), 5. 08 (1H, m), 4. 33 (1H, m), 3. 0 0 (4H, m), 2. 50-1. 10 (10H, m)

Purity		Example No.
91% (HPI		40.
.c)	6	. 89
		1H NER(6) ppm
٠		ppa
	Purity 91% (HPLC)	

WS	Purity > 6	Ž.	Example No.
503 (H+1)	>90% (NMR)		88
		300Ms, IMSO-d8 8. 31(IH, s), 8. 26and8. 06(2 H. MBq, J ^{eq} , 98ks), 7. 73and7, 22(4H, A'B' q, J ^{eq} , 77kz), 7. 5 0-7, 36(8H, m), 5. 10(2H, s), 4. 37(IH, tr.t., J ^e 12, 2Hz), 2. 38-2. 28(2H, m), 2. 10-1. 80(4H, m), 1. 70-1. 66(1H, m), 1. 50-1. 20(3H, m)	IH NAR(6) ppis

EP 1 162 196 A1

Table 20

Purity 約90% (NMR)	2	
MS 455 (0+1)		
Example No.	92 11	IH NAR(8) ppm
0 2001	7-3	300MHz, DMSO-d6 11.8(1H, brs), 8.07(1H, s), 7 80(1H d Tes 7Hz) 7 84(
	J 	z), 7. 69 (2H, , 4. 42 (2H, s)
0		(4H, m), 2, 40-1, 40 (10H, m)
Purity >90% (NMR)	<i>R</i>)	
MC AIG(N+I)		

Purity K

>90% (NMR) 537 (H+1)

	455 (M+1)	SF
	ty 約90% (NMR)	Purity
8.31(H, s). 8.27(H, d) [=8 .7Hz), 8.06=8.03(H, m), 7. 77-7.58(6H, m), 7.31(2H, d, J=8.7Hz), 5.81(2H, s), 4.40 (IH, m), 2.50-1.20(10H, m).	Constant of the constant of th	, =-•
300MHz, DASO-d6		
IH NAR(8) ppm	Example No. 91	Exam

İ	Purity >90% (NMR)		Example No.	MD 434 (M+1)		Purity >90% (NMR)		Example No.
			96		l		\lor	28
		B. 31(1H, d., J=1. 3Ha), 8. 27(1H, d., J=8. 8Hz), 8. 05(1H, d., 1H, d., J=8. 8Hz), 8. 05(1H, d., 1-8. 8Hz), 7. 76(2H, d., J=8. 7 1Hz), 7. 40-7. 25(4H, m), 7. 06 -6. 90(3H, m), 4. 53-4. 26(5H -8.), 2. 40-2. 18(2H, m), 2. 12 -1. 56(6H, m), 1. 50-1. 19(3H	IH NMR(8) ppm				300Hs, DMSO—d8 12, 9(1H, brs), 8, 02 (1H, s), 7, 82 (2H, m), 7, 40-7, 25 (5H, m), 4, 58 (2H, s), 4, 09 (1H, m), 3, 71 (1H, m), 3, 49 (2H, m), 3 21 (2H, m), 2, 35-1, 30 (14H, m),	1H NAR(8) ppm

ä

õ

Example No.

94

1H NAR(8) ppm

300MHz, DMSO-d6 8, 32(1H, a), 8, 27(1H, d, J=9 .0hz), 8, 05(1H, d, J=8, 7Hz), 7, 76-7, 70(3H, m), 7, 56(1H ,d, J=8, 4Hz), 7, 55-7, 35(6H ,a), 7, 22(2H, d, J=8, 7Hz), 5, .11(2H, s), 4, 36(1H, m), 2, 4 -2, 15(2H, s), 4, 36(1H, m), 2, 7 H, m), 1, 95-1, 75(2H, m), 1, 7 5-1, 55(1H, m), 1, 55-1, 20(3 H, m)

EP 1 162 196 A1

Purity

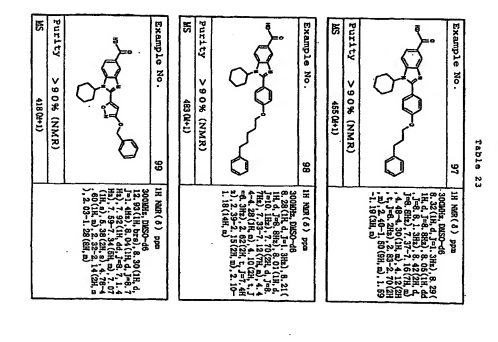
>90% (NMR) 531 (9+1)

300£iz, DaSO-d6
8. 32(1H, a), 8. 22(1H, d, J=8, THz)
8. 32(1H, a), 8. 22(1H, d, J=8, THz)
7. 72(2H, d, J=8, THz), 7. 31(4H, t)
7. 73(3H, a), 7. 21-7. 17(4H, m)
7. 73(3H, a), 4. 26(1H, t.)
7. 31(3H, a), 4. 26(1H, t.)
7. 31(3H, a), 4. 26(1H, t.)
7. 31(3H, a), 2. 50-2. 20(2H, a), 2. 50-2. 20(2H, a), 2. 50-2. 55(1H, a), 1. 75-1. 55(1H, a), 1. 75-1. 55(1H, a), 1. 155-1. 20(3H, a).

50

87

Example No.



ô.

g

8

t,

Example No. 102 IH.NuR(6) ppm Example No. 102 IH.NuR(6) ppm 200Hb, Das 5Hb), 7. 36(1H, a), 1. 86(1H, a), 1. 86(1H, a), 1. 86(1H, a), 1. 87(1H, a), 1. 87(1H, a), 1. 88(1H, a), 1. 88(1	11. NMR (6) 300MHz, DMS 12. 88 (1H, d, J 86 (1H, d, J H, d, J=8, J= (4H, AB9, J= 35 (5H, B), 5 (1H, B), 2. 3	Purity >90% (NMR)
--	---	-------------------

E	Ā	2	[F]
	Purity		Example No.
ŀ	9 <	A.	No.
427 (M+1)	>90% (NMR)		
Ė	NMR)	Ţ	
			8
	Ü	300MHz 8.46(1 1H, 8), 1Hz), 7.68(-7.30(8.5Hz), 98(1H), 2.00(1H)	H NE
		B 5H 88	1H NMR(8) ppm
		5-1. 11. 15. 15. 15. 15. 15. 15. 15. 15. 1	ē
		8. 16(8. 5, 2 8. 5+2) 7. 7. 55 4. 4. J= 4. 25-4 8. (2H, B	

8

EP 1 182 196 A1

ı		. 4	Ę
619 (H+1)	>90% (NMR)		Example No. 1
	(3H, p)	H. ABq. J=6 (Hz), 7. (48and; 87(2) H. ABq. J=6 (Hz), 7. (48and; 7. 4 6-7. 33 (6H, m), 6. 95and6. 76 (2H, 18 q, J=8, 2Hz), 6. 82 (1H, s), 5. 13 (2H, s), 4. 30 (1H brt. J=1. 2Hz), 2. 39-2. 18 (2H, m), 1. 98-1. 7(4H, m), 1. 48-1. 20	105 IH NUR(6) ppm 300MHz, DUSO-d6

	519 0(+1)	8
-	Pur1ty >90% (NMR)	u z
), 1. 95-1. 77 (4H, m), 1. 66-1 .56 (1H, m), 1. 46-1. 10 (3H, m	0	
J=8.7Hz), 7.32-7.09(9H, m) , 5.13(2H, s), 4.28(1H, brt, T=12.7Hz), 2.36-2.19(2H m		ŧ
12. 75 (1H, s), 8. 23 (1H, s), 7 .94and7. 86 (2H, ABq, J=8, 6H z), 7. 64and7. 05 (4H, A' B' q,		1
1H NMR(5) ppm 300MHz, DMSO-d6	Example No. 104	X

אכ	Purity		Example No.
401 (1+L)	>90% (NMR)		lo.
•	શ	V	103
		300Mis, DMSO-46 12. 79(1R, br.s), 8, 22 (2H, s) 12. 79(1R, br.s), 8, 22 (2H, s) 8, 02-7, 78(4H, s), 7, 63-7, 42 (6H, s), 7, 20-7, 09 (2H, s) 4, 43 (2H, s), 4, 27 (1H, br.t, 1-12, 22h ₂), 3, 69 (2H, s), 2, 9-2, 15 (2H, s), 1, 98-1, 72 (4 H, s), 1, 68-1, 59 (1H, s), 1, 4 3-1, 12 (3H, s)	IH NER(6) ppm

>90% (NMR)

429 (H+1)

¥, Purity

Example No.

107

1H NUR (8) - ppm

300MHz, DASO-46
12, 98 (H, brs.) 9, 92 (IH, brs.) 8, 93 (H, brs.) 7, 74 (M, br

Table 26

106

EP 1 162 196 A1

Example No.

Purity Example No. >90% (NMR) 108 JH NAR(6) ppm

3004hb, DASO-d6
8.24(1H, s), & Oland7, 90(2
R, ABo, J=6, Ths), 7, 65and7,
03(4H, A' B' q, J=8, Ths), 7, 31
2-7, 20(3H, m', 7, 08-7, 03(1
H, m), 4, 32(1H, brt., 1-12, 2H
st), 3, 77(3H, s), 2, 36-2, 20(2H, m), 2, 00-1, 78(4H, m), 1,
71-1, 59(1H, m), 1, 44-1, 11(3H, m) Purity

>90% (NMR)

429 (H+1)

S

443 (H+1)

Purity

5 497 Q(+1)

8

>90% (NMR)

112

Example No.

Purity Example No. >90% (NMR) 499 (M+1) 114 IH NAR (8) pps

8

Purity E

>90% (NMR)

471 (H+1)

93

>90% (NMR)

497 (H+1)

ä

Example No.

111

1H NAR(8) ppm

ŧ

¥ Purity

ĸ

ä

300Hiz, DMS0-d6 8. 22(1H, a), 7. 91 and 7. 87(2 H, M84, J=8, 71kz), 7. 68and 7. 18(4H, A' B' q, J=8, 71kz), 7. 3 6(1H, t, J=8, 51kz), 6. 80(1H, d, J=9, 01kz), 6. 72=6, 68(2M, d), 4. 30(1H, brt., J=12, 21kz), 3. 94(2H, t, J=6, 51kz), 2. 39 -2. 18(2H, m), 1. 97-1, 58(7H, m), 1. 45-1, 20(3H, m), 0. 97 (3H, t, J=7, 4Hz)

5 Purity

>90% (NMR)

471 (3+1)

300/Hts, DMSD-d6 12, 78 (1H, s), 8, 23 (1H, s), 7 12, 78 (1H, s), 8, 23 (1H, s) 9, 95 mod 7, 86 (2H, Ma, J=8, 9H s), 7, 89 mod 7, 18 (4H, A' B' q, J=8, 94, 25, 15 (1H, t, J=8, 3) Hs), 6, 81=4, 65 (3H, s), 6, 41 (2H, brs), 4, 54 (2H, d, J=8, 6) Hs), 4, 31 (1H, brt, J=12, 2Hs), 2, 41=2, 18 (2H, m), 1, 99=1 , 76 (4H, m), 1, 73 (3H, s), 1, 7 0-1, 58 (1H, s), 1, 68 (3H, s), 1, 45=1, 17 (3H, m)

IH NAR (&) ppm

Example

No.

113

1H NAR(6) ppm

300Mis, DMS0-d6
12. 73(1H, s), 8. 22(1H, s), 7
12. 434nd7. 85(2H, AB, J-9. 3H
15. 7. 61and7. 01(4H, K' B' q,
J-8. 6Hs), 7. 25-7. 00(4H, m)
J-8. 6Hs), 7. 25-7. 00(4H, m)
J-8. 6Hs), 4. 29(1H, brt., J-1,
J-8. 6Hs), 4. 29(1H, brt., J-1,
J-8. 6Hs), 4. 29(1H, brt., J-1,
J-8. 6Hs), 2. 38-2. 18(2H, m), 1
2. 2Hs), 2. 38-2. 18(2H, m), 1
3. 2Hs), 1. 61(3H, m), 1. 60(3H, m)
Hs), 1. 48-1. 16(3H, m)

Table 28

EP 1 162 196 A1

Table 29

Example No. 117 1H NAR(6) ppm 300MHs, DMSO-d6 12.8(1H, brs), B. 1.8(1H, br
117 IH NAR(6) p 300MHz, DMSO- 12. 8(1H, by) 7. 98(1H, d., J=8, 8H) 17. 98(2H, d., J=8, 8H) 17. 58(2H, d., J=8, 11, d., J=8, 2H) 17. 51(2H, d., J=8, 11, d., d., d., d., d., d., d., d., d., d.
1H NAR (6) ppm 3002Hz, DASO-d6 12. 8 (1H, brs), d. Je. 17. 98 (1H, d. Je. 18. 18. 19.
55.27 55
22(1H, 9) 71, 71, 87 71, 80(2H, 0, 1) 71, 67(3H, 1) 71, 71, 71, 11 12, 7, 41 12, 7, 41 12, 7, 41 12, 7, 41 12, 7, 41 12, 7, 41 12, 7, 41 13, 7, 7, 7, 7, 7, 7, 7, 7, 7, 7, 7, 7, 7,
9889E8

Purity SW

>90% (NMR)

300MHz, DMSO-d6
8. 23(1H, s), 7, 99 (1H, d, J=8, 4Hx)
7. 61smd7, 18 (4H, Mbq, J=8, 6Hz), 7, 30-7, 22 (2H, m), 7, 0
1 (2H, d, J=6, 1Hz), 6, 92 (1H, m), 4, 28 (1H, m), 4, 25 (2H, t, J=7, 2Hx), 3, 28 (3H, m), 4, 28 (2H, t, J=7, 2Hx), 2, 38 (3H, m), 2, 1, 70-1, 55 (1H, m), 1, 15 (3H, m)
1, 10 (3H, m), 1, 50-1, 15 (3H, m), 1, 15 (3H, m)

471 (01+1)

å

Example No.

120

IH NMR(&) ppm

g

- N.	MS 567 (H+1)	Purity >80% (NMR)		Example No.
	H-1)	(NMR)	ÿ	116
		(1H, m), 1.86-1.20(3H, m).	300,HH, DISO-d6 8. 30(1H, d.) F-8. 8Hz) 9Hz) E. 03(1H, d.) F-8. 8Hz) 7. 68(2H, d.) F-8. 8Hz) 7. 24 (2H, d.) F-7. 2Hz) 7. 19-7. 10 (6H, m.) 6. 94(2H, t.) F-7. 2Hz (6H, m.) 6. 94(2H, t.) F-7. 2Hz 1. 4. 34(1H, m.) 4. 19(4H, brs 1. 4. 34(1H, m.) 4. 19(4H, brs 1. 4. 34(1H, m.) 4. 17(1-1), 55 (2H, m.) 2. 10-1, 15(2H, m.) 1. 75-1, 75(2H, m.) 1. 75-1, 55	IH NMR(6) ppm

MS 499 (N+1)	Purity >90% (NMR)	9-9-20-	Example No.
	R)	300MHz, DKS0-d8 8.23 (IH, s) 7.9 8.48c, J=8.68b2) 19 (4H, A' B' c, J=6 5 (IH, t, J=7.8Hz) 9 (3H, b) 4.30 (17 2 Hz) 4.00 (2H, J=7.20 (2Hz) 5, 0.93 (6H, d, J=6	115 IH NMR(8) ppm
		7. 93and7. 87(2 12), 7. 69and7. 13), 7. 69and7. 13H2), 6. 82-6. 6 10(114, brt, J=1. 214, t, J=6. 94s) 224, t, J=6. 94s) 17-1. 20(34, a)	

MS 671	671 (H+1)	
 Example No.	119	1H NMR(6) ppm
} } !		74and6, 90(4
\{\bar{\}}	£0)	.31(21, t.]-6, 8Hz) 3, 74(3, 1, 2), 3, 74(3, 1, 2), 3, 74(2H, t.), 2, 196, 7Hz), 2, 30(2H, m), 2, 02(2H, m), 1, 55 86(2H, m), 1, 63(1H, m), 1, 55 -1, 15(2H, m)
Purity >90%	>90% (NMR)	
HS 471	471 (8+1)	

Purity Example No. >90% (NMR) Table 30 118 300MHz, DMSO-d6 13. 31H, brs), 8. 30(IH, s), 8. 25 (IH, d, J=8, 9Hz), 8. 04(IH, d, J=8, 7Hz), 7. 72(ZH, d, J=8, 8Hz), 7. 75 (MH, d, J=8, 6 Hz), 7. 47 (4H, d, J=8, 6Hz), 7 13. (ZH, d, J=8, 9Hz), 6. 84(1 IH, s), 4. 33 (IH, m), 2. 45-2. 1 0(ZH, m), 2. 10-1. 95 (ZH, m), 1. 95-1. 70 (ZH, m), 1. 70-1. 5 5 (IH, m), 1. 55-1. 15 (3H, m). 1H NAR(8) ppm

EP 1 162 198 A1

ŧ

5 Purity

>90% (NMR)

517 (H+1)

Example No.

126

1H NMR(&) ppm

300NHz, DMSO-d6
8, 32(1H, s), 8, 14(1H, d,]=8
7Hz), 8, 03(1H, d,]=6, 7Hz)
7, 77(2H, d,]=9, 0Hz), 7, 52
-7, 31(7H, m), 6, 74 (2H, m), 5
-26 (2H, m), 4, 61 (1H, m), 2, 9
6 (1H, m), 2, 60-2, 10 (5H, m).

MS . 441 (H+1)	Purity > 90% (NMR)		Example No.
			122
		000Miz. DMSO-d6 12.8 (1H. brs), 8. 22 (1H. s), 7. 87 (2H. m), 7. 62 (2H. d) 1=8 .1Hz), 7. 60-7. 20 (7H. m), 5. 23 (2H. s), 4. 46 (1H. m), 2. 50 -2. 30 (2H. m), 1. 70-1. 40 (10 H. m).	IH NAR(6) ppm

		SK
	Purity > 90% (NMR)	ğ
3.76(3H, s), 3.07(2H, t, 12-6 .712), 2.29(2H, m), 2.00-1, 75(4H, m), 1.70-1, 55(1H, m)		
.17 (4H, ABq, J=8. 7Hz), 7.24 (1H, m), 6.94 (2H, m), 6.82 (1		
12. 85 (1H, brs), 8. 24 (1H, s) , 8. 01 (1H, d, J=8. 7Hz), 7. 90 (1H, d, J=8. 6Hz), 7. 62and, 7		é
300MHz, DMSO-d6		
1H NUR(8) ppm	Example No. 121	EX

G

Example No.

125

IH NAR(8) ppm

300MHs, DMSO-d6 8.32(H, s), 8.28(H, d, J=8 7.12), 8.05(H, d, J=9.0Hz), 7.43 (4H, d, J=9.0Hz), 7.36-7.20 (8H, m), 4.74(2H, d, J=7.5Hz), 4.3 8(H, m), 4.74(2H, d, J=7.5Hz), 4.3 8(H, m), 2.40-2.15(2H, m), 2.40-2.15(2H, m), 2.40-2.15(2H, m), 1.95-1.8 5(2H, m), 1.85-1.55((H, m), 1.55-1.20(3H, m), 1.55-1.20(3H, m).

533 (H+1)

Purity >90% (NMR)

3000Hz, DMSO-d8
13.1(H, br.s), 8.29(H, s),
8.17(H, d.J-8, Ttz), 7.99(
H, d.J-8, Ttz), 7.77(2H, d.J-8, Ttz), 7.77(2H, d.J-8, Ttz), 7.77(2H, d.J-8, Ttz), 6.76
6.72(2H, d.J-8, 3H, 3), 6.76
6.72(2H, d.J-8, 3H, 3), 6.76
6.72(2H, d.J-8, 3H, 3), 6.76
1.72(2H, d.J-8, 3H, 3), 6.76
1.72(2H, d.J-8, 3H, 3), 6.76
1.72(2H, d.J-8, 3H, 3), 6.76
1.73(2H, d.J-8, 3H, 3), 6.76
1.75(2H, d.J-1, 7.75-1, 56(1H, d.J-1

124

1H NAR(&) ppm

Table 32

EP 1 162 186 A1

Example No.

8

Purity

>90% (NMR)

425 (H+1)

MS 505 (M+1)	Purity >90% (NMR)	C		=0	Example No.
	-1.69(1H, D), 1.44-1.19(3H	J.R. 62, 243, 6, 75 (JH, 6), 4, 36 J.R. 243, 6, 75 (JH, 6), 4, 36 -4, 18 (JH, m), 2, 38-2, 17 (2H , m), 1, 95-1, 76 (4H, m), 1, 70	22(4H, A'B q, J=8, 6Hz), 7.5 22(4H, M'B q, J=8, 6Hz), 7.5 2-7.39(1H, m), 7.47and7.41 (2H, A'B q, J=8, 1Hz), 6, 91(3000Hz, DMSO-d6 8. 21 (1H, s), 7. 92end7. 86 (2	129 IH NAR(8) ppm

nd7.85(.61and7. 61a, 7.; 4(14, 5r. 2.15(2H 2.1.70

ಕ

3

127 IH NAR(&) DPM 300MHz, DMSO-46 13 2(IH, brs), 8, 33(IH, s), 8, 12(IH, d, J=8, 718), 7, 78(2H, d) 14 (J=8, 81s), 7, 78(2H, d) 15-8, 718), 7, 62-7, 32(7H, m) 2, 25(2H, m), 2, 25-1, 50 (6H, d) 2), 35(2H, m), 2, 25-1, 50 (6H, m)	Purity >90% (NMR)		Example No.
		5 (2) (2) (2) (2) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4	

ü

8

>90% (NMR) 5900H1)

õ

Example No.

Table 34

130

1H NAR(6) DDM

-300HHs, DMSO-d6

8. 27(H, s), 7. 69(2H, d, J=8

6. 17, 49-7, 21(11H, m), 5

-08and6, 03(2R, A8q, J=12.6

Hs), 8. 607-4. 99(1R, m), 4. 26

(2H, d, J=0. 6Hs), 2. 40-2. 18

(2H, d), 2. 04-1.77(4H, m), 1

.70-1. 68(1H, m), 1. 48-1. 15

(3H, m)

EP 1 162 196 A1

Purity

B

Example No.

1H NACR (&) ppm

131

300Mis, DMS0-d6 8. 29(1H, a), 8. 11 (1H, d, J=9 .0Hs), 7. 96(1H, d, J=8, 4Hz) 7. 80(2H, d, J=8, 1Hs), 7. 72 -7. 41 (7H, m), 7. 12 (1H, d, J= 12. 6Hs), 7. 01 (1H, d, J=8, 4H a), 6. 12 (2H, s), 4. 08 (1H, m) .2. 15-2, 10 (2H, a), 2. 00-1, .75 (4H, m), 1. 75-1, 55 (1H, m) .1. 60-1, 20 (3H, m)

Example No. Purity ĸ >90% (NMR)

589 (N+1)

132 1H NAR(6) ppm

300Δis, DuSO-d6 12. 8 (IH, brs), 8. 23 (IH, s), 7. 97 (IH, d, J-8, Thz), 7. 87 (IH, d, J-8, 6Hz), 7. 66 (2H, d, J-8, 6Hz), 7. 49-7, 33 (FH, s), 7. 17-7. 05 (6H, s), 5. 12 (2H, s), 4. 31 (IH, s), 2. 40-2. 15 (2H, s), 2. 05-1. 20 (6H, s).

>90% (NMR)

Purity Z

519 (J+1)

ğ

1/2

	MS · 531 (H+1)
	Purity .>90% (NMR)
8H, 8), 2, 03-1, 84 (4H, m), 1, 1, 17 (1H, m), 1, 46-1, 20 (3H, m)	7
, J=8.0Ha), 7.09(2H, d, J=8. 7Ha), 6.28(1H, s), 4.37(1H,	
7.61(2H, d, J=8, 7H2) 7.61(2H, d, J=8, 7H2), 7.31 (4H, d, T=8, 0H2), 7.16(4H, d	
3002Hz, DASO-46 8. 57(1H, s), 8. 01(1H, d, J=8	
IH NAR(6) ppm	Example No. 133

EP 1 162 186 A1

Table 36

EP 1 162 196 A1

> 9 0% (NMR) 547(H+1)

Purity MS

	Example No.
a, o	138
3001/H, DISO-d6 12,776(1H, brs), 8, 22(1H, s) 7,76(1H, d, J=8,7Hz), 7, 85 (1H, d, J=8,7Hz), 7, 54-7, 49 (4H, m), 7, 42-7, 21(5H, m), 7, 11-7, 09(3H, m), 6, 39(1H, m)), 6, 17(2H, s), 4, 29(3H, m), 3, 11(2H, m), 2, 40-2, 20(2H, m), 1, 99-1, 23(8H, m)	1H NAR(6) ppm

	469 (H+1)	æ
	ity >90% (NMR)	Purity
300111, DISO-46 8. 24(1H, s), 8. 11(1H, s), 7. 76(2H, d, J=9, 0Hs), 7. 37-7, 18(7H, m), 4. 43-4, 30(1H, m), 4. 13(2H, t, J=6, 3Hs), 2. 84 -2. 88(5H, m), 2. 42-2. 22(2H m), 2. 18-1. 80(6H, m), 1. 70 -1. 20(4H, m)		ž 💆
TH MAR(&) DDD	Example No. 137	ž.

MS 471 (M+1)	Purity > 90% (NMR)		Example No.
		0	136
		300HH, DUSO-d6 8, 13(1H, 9), 7, 55(2H, d,]-6 7ha), 7, 63(1H, e), 7, 35-7, 12(7H, m), 4, 35-4, 20(1H, m), 4, 10(1H, t,]-6, 312), 2, 78 (2H, t,]-7, 5Hs), 2, 33-1, 78 (8H, m), 1, 70-1, 16(4H, m)	IH MAR(8) ppm

ន្ត

12.83(2H. brs.) 8.22(1H. a) (7.14 (4.17 + 2.18 - 71h.) 7.84 (7.18 (4.17 + 2.18 - 71h.) 7.85 (7.18 (4.17 + 7.63 - 7.63 - 7.63 - 7.63 (1h. a), 7.85 (7.18 (4.17 + 7.63 - 7.63 (1h. a), 7.85 (1h. a),
--

	MS 647 (M+1)
	Purity >90% (NMR)
300MHs, DMSO-46 12. 73 (1H, brs), 8. 22(1H, s) 7. 93 (1H, d, J=8. 7Hz), 7. 73 (1H, m), 7. 60-7. 57 (2H, m), 7. 47-6. 90 (1H, m), 5. 11 (2H, s) 14. 33-4. 28 (3H, m), 3. 09-3 04 (2H, t, J=6. 7Hz), 2. 35-2 20 (2H, m), 1. 95-1. 10 (8H, m)	ord of
IH NUR(8) ppm	Example No. 139

EP 1 162 196 A1

EP 1 162 198 A1

Table 38

8		8	Ġ	8		4	g	2	,	3	ā i	š
SW	Purity			Example	MS MS		<u> </u>	Example	SW	Purity	<u>.</u> .	Example
585 (N+1)	>90% (NMR)	0	to, A	No. 144	>90% (NMR) 867(H+1)	2		No. 143		>90% (NMR)		No. 142
	. 10 (3H, m).	1, 2, 5, 08 (2H, 8), 4. 1, 2, 40-2, 15 (2H, m), 2, 05-1, 70 (4H, m), 2, 05-1, 70 (4H, m), 1, 50	300HHz, JMSO-G6 13, 0(1H, brs), 8, 31(1H, s), 13, 0(1H, d., J=8, 7Hz), 8, 04(8, 23(1H, d., J=8, 7Hz), 7, 80(2H, d. 1H, d., J=8, 7Hz), 7, 76(3H, m) 1, 56-7, 40(4H, m), 7, 03-8	1H NAR(8) P	(a), 1.55-1.15(3H, b)	=8.842), 7.01(24, d) =8.842), 7.01(24, d) z), 5.11(24, s), 4.35 3.79(34, s), 2.45-2 m), 2.15-1.95(24, s)	33. 1(IH, brs), 8. 30 (IH, s), 13. 1(IH, brs), 8. 30 (IH, s), 8. 24 (IH, d, J=8, 8Hz), 8. 03 (IH, d, J=8, 7Hz), 7. 74-7. 71 (IH, d, J=8, 3Hz), 7. 40-7. 36 (3H, d, J=8, 3Hz),				7Hz), 8, 05(1H, d) Jei, 7, 76-7, 72(3H, a), 7, 76-7, 72(3H, a), 7, 39-7, a), 5, 11(1H, a), 4, 31, 2, 35(3H, a), 2, 35-2, 35(2H, a), 2, 35(2H, a), 1, 75-1, 75(2H, a), 1, 55-1, 15(3H, a)	1H NMR (6 300MHz, D1 8, 32 (1H, 1

ğ

ğ

Example No. 147 1H NAR(8) ppm S

555 (H+1)

300MHz, CDC13
8.61(H, s), 8.04(1H, d, J=8, 7Hz)
7.66(1H, d, J=2, 4Hz), 7.59
(2H, d, J=8, 7Hz), 7.42(1H, d, J=8, 2Hz), 7.89
(2H, d, J=8, 7Hz), 7.28(2H, d, J=1, 8Hz), 7.28(2H, d, J=1, 8Hz), 7.28(2H, d, J=1, 8Hz), 7.28(2H, d, J=1, 8Hz), 7.28(2H, d, J=1, 2Hz), 7.03(2H, d, J=8, 7Hz), 4.94
(2H, s), 4.37(1H, m), 2.43-2.
21(2H, m), 2,17-1.88(4H, m), 1.79(1H, m), 1.43-1.28(3H, m), 1.

Purity

>90% (NMR)

553 Q+1)

Purity Example No. >90% (NMR) 146

1H NMR(&) ppm

8

똢 Purity

>90% (NMR)

593 (J+1)

300Miz, DMS0-d6 8.31(H, s), 8.23(H, d, J=8, His) 8.812, 8.02(H, d, J=8, His) 7.73-7.71(3H, m), 7.54(H d, J=8, 2Hz), 7.48(2H, d, J= 8.4Hz), 7.41-7.37(3H, m), 7. 7.27(2H, d, J=8, His), 7. 1.27(2H, d, J=8, His), 5.13(2 H, s), 4.34(H, m), 2.40-2.2 1.96-1.76(2H, m), 1.70-1.6 5(H, m), 1.50-1.15(3H, m), 1.31(9H, s).

145 IH NAR(8) ppm

Table 39

Example No.

EP 1 162 198 A1

ä

S Purity

>90% (NMR)

587 (1+1)

Example No. 150

300MHz, DMSO-d8 8. 22 (IH, 6), 8. 85 (IH, d, J=8 4Hz), 7. 88 (IH, d, J=8, 7Hz) 7. 65 (IH, d, J=8, 4Hz), 7. 52 -7. 28 (7H, a), 7. 23 (2H, d, J= 9. 3Hz), 7. 14 (2H, d, J=8, 7Hz)), 6. 14 (2H, d), 3. 90-3, 72 (IH, d), 2. 20-1, 10 (10H, m)

IH NAR(8) ppm

300Mbs, DMSO—d6
8. 24 (1H, d.) —1. 6Hs), 7. 96 (
1H, d.) —9. 0Hs), 7. 88 (1H, dd
.] —9. 0, 1. 5Ms), 7. 88 (1H, d,
.] —9. 7. 12. 7. 50 –7. 30 (6H, m)
.7. 22 –7. 00 (6H, m), 5. 13 (2H
., b), 3. 98 –3. 80 (1H, s), 2. 36
—1. 10 (10H, m)

IH NAR(6) ppm

149

Purity

>90% (NMR)

557 (X+1)

ጅ

Example No.

1H NAR(6) PRIM
300MHz, DMSO-46
8, 21 (s. 1H), 7.89 (1H, d. J=8, 7Hz), 7.87 (1H, d. J=8, 7Hz)
7, 163, 7.46 (5H, d.), 7.30-7, 12 (5H, d.), 7.08 (1H, d., J=1, 1, 0Hz), 8, 81 (1H, d.), 8, 12 (1H, d.), 2, 12 (4H, d.), 2, 12-2, 06 (2H, d.), 1, 89-172 (4H, d.), 2, 12-2, 06 (2H, d.), 1, 40 (2H, d.), 2, 12 (4H, d.), 2, 12 (

Table 148

Example No.

EP 1 162 186 A1

	489 (M+1)	፳
	>90% (NMR)	Purity
1. 45-1. 15 (3H, n)		
z), 4. 20 (1H, brt, J=12. 2Hz) , 2. 32-2. 13 (2H, m), 1. 92-1.		<u></u>
8. 20(1H, s), 8. 93and7. 83(2 H, ABq, J=8. 7Hz), 7. 86-7. 21 (11H m) 7. 03(2H d 1=8. 7H		<u>_</u>
300Miz, DMSO-d6		
IH NAR(&) ppm	No. 153	Example No.

Ġ

ŧ

SW	Purity . >9		Example No.
456 (M+1)	>90% (NMR)		152
		0. 21 (47, m), 1. 99 7, 20 (27, m), 7, 63 -7, 20 (91, m), 4, 20 - 15 (21, m), 1. 95 -1, 74 (41, m), 1. 70 - 1, 54 (11, m), 1. 44 -1, 14 (31, m)	56 9

ŧ

8

G

8

ä

ö

Purity Example No. >90% (NMR) 605 (H+1) 151 1H NMR (5) ppm

300MFz, DMSO-d6

8. 18 (14, s) 7, 78-7, 78 (3H, s) 7, 78 (3H, s) 7, 78 (3H, s) 7, 70 (12H, d, jed, 71x) 4, 48 (1H, d, jed, 71x) 4, 88 (1H, d, jed, 71x) 4, 22 (1H, d, jed, 71x) 4, 23 (2H, s) 2, 37-2, 16 (2H, s) 1, 19 6 -1, 76 (4H, s) 1, 1, 64 (1H, s) 1, 1, 48-1, 14 (3H, s)

Table 41

EP 1 162 196 A1

EP 1 162 196 A1

6		6	4	6	G	6	ii.		6		u	•	•
MS.	Pu		₹ ·	텧	: 12 12 12 13		3	E	ES.	P			F
	Purity	O'	ک ک	Example No	MS MS		<u> </u>	Example No		Purity	. (Example No
626 (4+1)	1) %08<	Ç	Ţ		> 9 0 % (NI	C	¢ l		489 (H+1)	>80%	C	5	
5	(NMR)) ا	ļ Į	156	(NMR)	٠, C	, י	155	ž	(NMR)			154
	03(5H, m), 1. 39(9H, s)	. 30(1H, brt, J=12. . 30(1H, brt, J=12. 9(2H, brd, J=12.0H, brd, J=6.3Hz), 2. (2H, d, J=6.3Hz), 2. (2H, m), 2.38-2.20(HZ, JMSO-do (1H, s), 7. 92an (q, J=8, 7Hz), 7. H, A' B' q, J=8, 7 t, J=8, 5Hz), 6	1H NAR(6) ppm	8), 1. 00-0. 82 (2H, m)	10, 21, 27, 28, 28, 21, 21, 21, 21, 21, 21, 21, 21, 21, 21	, DISO-46 H, 8), 7.85 J=8, 7Hz), A, B, Q, J=8	1H NAR(8) ppm			brt, J=12. 2Hz), 2. 37-2. 18(2H, m), 1. 98-1. 77 (4H, m), 1. 70-1. 58 (1H, m), 1. 48-1. 20(3H, m)	8. 23 (1H, s); 7. 94snd7. 86(2 H, ABq,]=8. 6Hz); 7. 72-7. 16 (13H, m), 5. 25 (2H, brs), 4. 5 5 (2H, d, J=8. 6Hz); 4. 31 (1H,	TH NAR(

Table 42

É

	MS 517 (M+1)
	Purity >90% (NMR)
308HJa, DMSO-d8 12.76 (IH, brs), 8. 22 (1H, s) 7. 93 (2H, d. J=8. THz), 7. 85 (2H, d. J=8. 6Hz), 7. 53-7. 21 (10H, m), 6. 94 (2H, d. J=8. TH z), 4. 30-4. 12 (3H, m), 3. 05 (2H, m), 2. 35-2. 15 (2H, m), 1. 95-1.75 (4H, m), 1. 75-1. 55 (1H, m), 1. 50-1. 10 (3H, m)	
IH NAR(8) ppm	Example No. 158

l:

g

ધ

ŧ

L	J.	627 (H+1)	SW
8 -	(2H, m), 1, 99-1, 79 (4H, m), 65 (1H, s), 1, 49-1, 15 (3H,	/ >90% (NMR) .	Purity
77.5.6.4.88	12.78 (1H, bars), 8.22 (1H, 12.78 (1H, bars), 8.22 (1H, 1.7.86 (1H, d, J=8.6Hz), 7.8 (1H, d, J=8.6Hz), 7.60 (2H, d, J=8.2Hz), 7.60 (2H, dd, J=8.3Hz), 7.48 (1H, d, J=8.3Hz), 7.48 (1H, d, J=8.3Hz), 7.48 (1H, d, J=8.3Hz), 7.18 (2H, d, J=8.4Hz), 6.73 (2H, s), 5.08 (2H, s), 4.23 (1m), 3.69 (9H, s), 2.37-2.1 (m), 3.69 (9H, s), 2.37-2.1		8
	1H NHR(8) ppm	le No. 157	Example No.

EP 1 162 198 A1
Table 43

Table .

	Purity	•	Example No.	8	Purity	,	Example No.	1
498 (H+1)	>90% (NMR)		o. 162	526 (H+1)	>90% (NMR)		0. 161	0.0 (M · 5./
	-1. 11 (3H, m)	300Hfs, DMSO-46 12. 87(1H, brs), 8. 68(1H, d, J-6, 0Hz), 8. 23(1H, s), 7. 99 and 7. 80(2H, ABq, J-8, 6Hz), 7. 61and 7. 18 (4H, A Bq, J-8, 6Hz), 7. 61and 7. 18 (4H, A Bq, J-8, 11, 2H, ab), 6. 29(1H, brs), 4. 28(1H, brt, J-12, 24, 25, 27-2, 11(2H, ab), 6. 20-1, 71(4H, ab), 1. 92(1H, ac), 1. 46	TH NAK (9) DDD		6-1. 15(15H, B)	3.00 S(H) brs), 8. 76(1h, brs), 8. 05(H) brs), 8. 19and8, 00 (2H, A50, 1=8, 3Hz), 7. 79and8, 17. 25(4H, K) 8. 0, 1=8, 3Hz), 7. 25(4H, K) 8. 0, 1=8, 3Hz), 7. 39(H) brs), 6. 88-6. 73(4H, brs), 6. 88-6. 73(4H, brs), 6. 80-6. 73(4H, brs), 7. 80-6. 73(4H, brs), 80-6. 7	(6) p	-

Example Mo. 160 IH NGR(\$) ppm

300MHs, DIX50-d6
8.90(IH, brs), 8.59(Ih, brs)
8.80(IH, brs), 8.18end8.00
(2H, Alo, Jes, 5Hs), 7.7send
7.10(4H, A'B', Jes, 5Hs), 7.7send
7.10(4H,

EP 1 162 196 A1

300MHz, DMSO-d6 13.1(1H, brs), 8.32(1H, s), 8.28(1H, d, J=8, Hz), 8.05(1H, d, J=8, THz), 7.80-7.75(3H, m), 7.69(1H, d, J=4, 1Hz), 7.57(2H, m), 7.34-7.29(3H, m), 5.24(2H, s), 4.39(1H, m), 5.24(5-2), 20(2H, m), 2.20-1, 95(2H, m), 1.75-1, 55(1H, m), 1.55-1, 15(3H, m)

1H NMR(8) ppm

		4	164	
	(1H, m), 1. 43-1. 19(3H, m)	300MHz, DMSO-68 8. 20 (1H, 8), 7. 87 (2H, 8), 7. 8. 20 (1H, 8), 7. 87 (2H, 8), 7. 68and7, 18 (4H, ABG, J=8, 7Hz, 6), 7. 1 (1H, 6), 1=9, 4Hz), 6. 72 (1Hs, 6), 71 (1H, 6), 1=6, 8Hz), 4. 8 (2H, 9), 4. 29 (1H, brt. J=12 2 (2H, 9), 4. 10 (1H, 1]=6, 7Hz), 2. 39 2. 19 (2H, 1), 1, 97-1, 78 (4H)	IH NMR(8) ppm	
	t	8 8		
MS 615 (N+1)	Purity >80%		Example No.	
+1)	>90% (NMR)	.60%	167	

벊

>90% (NMR)

497 (N+1)

S Purity

Ġ

ž

Purity

â

Example No.

8

ď

Example No.

8

S

ü

Purity

>90% (NMR)

300MHz, DMSO-d6 8. 23 (H, s), 7. 55and7. 86 (2 H. ABa, J=6. 6Hz), 7. 69and7. 18 (4H, K B a, J=8. 6Hz), 7. 3 5 (HH, t, J=8. 6Hz), 6. 80 (H, d, J=7. 6Hz), 6. 72-6. 69 (2H, m), 5. 20 (HH, t, J=3. 7Hz), 4. 31 (HH, brt, J=12. 2Hz), 3. 95 (2H, t, J=6. 8Hz), 2. 49-2. 19 (4H, m), 1. 97-1. 76 (4H, m), 1 68 (3H, s), 1. 67-1. 64 (1H, m)), 1. 61 (3H, s), 1. 45-1. 20 (3 H. m)

611 (H+L)

5

Example No.

163

IH NMR(8) ppm

EP 1 162 196 A1

Table

õ Example No. Purity 돐 >90% (NMR) 583 (M+1) 166 300Miz, DMSO-d6 8. 27(1H, d.) 8. 13(1H, d.) 1=8 8. 27(1H, d.) 1=5, 0Hz) 7. 73(1H, d.) 1=1, 8Hz), 7. 68 (2H, d.) 1=6, 4Hz), 7. 54(1H, d.) d.) 1=8, 42, 1Hz), 7. 44(17, 31 (6H, m), 7. 19(2H, d.) 1=8, 4Hz 1, 5. 10(2H, s.) 4. 32(1H, m.) 2. 50(3H, s.) 2. 40-2. 16(2H, m.) 1, 55(1H, m.), 1. 55-1. 10(3H, m.) 1H NMR(8) ppm

167

1H NAR(6) ppm

300MHz, DMSO-d6 8. 25(IH, s), 8. 09(IH, d, J=8 4. 4b.), 8. 00(2M, d, J=8, 4Ms), 7. 94(1M, d, J=8, 7hz), 7. 80 (IH, d, J=2, IHz), 7. 73(2M, d, J=8, 1Lz), 7. 65(2H, d, J=8, 1.z), 7. 65(1H, d, J=8, 1.z), 7. 66(2H, d, J=8, 1.z), 7. 16(2H, d, J=8, 1.z), 5. 13(2H, s), 4. 30(IH, s), 3. 26(3H, s), 4. 30(IH, s), 3. 26(3H, s), 2. 40-1, 16(2H, s), 2. 5(3H, s), 2. 5(3H, s), 1. 75(4H, s), 3. 26(3H, s), 1. 55-1, 16(3H, s).

EP 1 162 198 A1

	MS 538 (X+1)
	Purity > 90% (NMR)
300Hz, DISO-46 300Hz, DISO-46 12.7 (1H, m), 8.66 (1H, s), 7. 8.61 (1H, m), 8.21 (1H, s), 7. 92-7. 79 (4H, m), 7. 61-7. 56 31, m), 7. 50-7. 43 (2H, m), 7. 10 (2H, d.), 78-71-35, 5.09 (2H, s), 4.26 (1H, m), 2.40-2.15 (2H, m), 2.00-1.75 (4H, m), 1. 76-1, 55 (1H, m), 1.50-1.15 (3H, m).	
IH NMR(8) ppm .	Example No. 170

(1+X) 129 SM	Purity >90% (NMR)		Example No.
	MR)		169
		300MH, DASO-d6 8 31(1H, p), 8.26(1H, d, J=8, TH±) . 712.) 8.05(1H, d, J=8, TH±) . 7.78-7, 71(3H, m), 7.59-7, 41(6H, m), 7.23(2H, d, J=9, 0 H2), 5.11(2H, s), 4.35(1H, m)), 2.40-2.15(2H, m), 2.15-1 . 15(2H, m), 1.95-1, 75(2H, m)), 1.75-1, 55(1H, m), 1.165-1 . 15(3H, m)	1H NMR(8) ppm

Table 47

EP 1 162 196 A1

MS 590 (M+1)	Purity >90% (NMR)		Example No. 174	Purity > 90% (NMR) MS 540(M+1)		Example No. 173	Purity > 90% (NMR) MS 537(M+1)		Example No. 172	
	1	300MHz, DESO-d6 12.80(1H, etc.), 6.26(1H, s) ,8.01(1H, d, J=8, 7Hz), 7.85 (1H, d, J=8, 7Hz), 7.80-7.70 (1H, m), 7.60-7.36(7H, m), 7.10-7.36 (1H, m), 7.60-7.36(7H, m), 2.32-1 ,4.11-3.80(1H, m), 2.32-1 ,18(14H, m)	9		8. 33(1H, s), 8. 29(1H, d, J=8 7Hs), 8. 66(1H, d, J=8, 7Hs), 7. 82-7. 74(4H, m), 7. 45(1H, d, J=8, 7Hs), 5. 28(2H, s), 4 dd, J=8, 43, 30, 32, 7. 39(2H, s), 4 40(1H, m), 2. 40-2. 15(2H, m), 2. 40-2. 15(2H, m), 1. 95-1 76(2H, m), 1. 76-1. 55(1H, m), 1. 55-1. 15(3H, m), 2. 40-2. 15(1H, m), 2. 40-2. 15(1	MAR(8) ppm DONHE, DKSO-d6		7. 88 (17. 88	300MHz DWSO-46	

EP 1 162 198 A1

S	Purity		example No	
568 (N+1)	>90% (NMR)		No.	
₹.	(NMR)	Y.	176	
	9(3H, s), 1.50-1.00(5H,	300MHz, DMSO-86 8.24(1H, 8), 7.97aind7.87; H, ASq. J=8.6Hz), 7.69and7. 19(4H, K'B'q, J=8.6Hz), 7.69and7. 19(4H, K'B'q, J=8.1Mz), 6.81(1); d. J=8.2Mz), 6.72(1H, 8), 6.72(1H, 8), 6.71(1H, 4), 3.95-3.75(3H, 3), 3.03(1H, 1, 3), 2.39-2.15(3H, 3), 2.39-2.15(3H, 3), 2.39-2.15(3H, 3), 2.39-2.15(3H, 3), 3.03(1H, 1, 3), 2.39-2.15(3H, 3), 3.95-3.75(3H, 3), 3.95-	1H NER(6) ppm	

SW	Purity		· ·		20	Example No	
(1+10) 868	>90% (NMR)	الم الم				o. 175	
	-J. 77 (28, 19)	-2. 16 (3H, m), 1. 96-1. 65 (6H -m), 1. 60-1, 13 (6H, m), 1. 10	7. 25 (2H, d, J=7. 7Hz), 5, 41 (2H, brs), 4. 54 (2H, d, J=6. 6	z), 7. 61and7.00(4H, A'B'q, J=8.5Hz), 7. 31=6.91(2H, m)	DMSO-d8 H, s), B. 21 (1H	TH NAR(6) ppm	

Table 49

EP 1 162 196 A1

EP 1 162 196 A1

8		8	å	8		
MS	Purity			Example No.	#S	,
536 (1+1)	>90% (NMR)			No. 180	457 (¥+1)	A CAN CHAIN
	66 (1H, m), 1. 49-1. 18 (3H, m)	6.95(1H, dd, J=9.05.19(2H, s).4.30.78(3H, s).2.40-2.78(3H, m).2.40-2.	8. 2(14, 8), (-91(14, d,]=8 .6Hz), 7. 85 (1H, d,]=8. 6Hz) ,7. 63 (2H, d,]=8. 4Hz), 7. 60 (1H, d,]=9. 0Hz), 7. 25 (2H, d ,1=8. 4Hz), 7. 23 (1H, d,]=3.	NAR (6) ppm OMPz, DMSO-d6		

MS 45	Purity >909		Example No.
457 (H+1)	>90% (NMR)	6	179
	2-1. 10(3H, m)	8. 32 (1H, a), 8. 29 (1H, d, J=9 (1H, d, J=9 (1H, d), 8. 60 (1H, d, J=3, THz) 7. 61 (1H, d, J=3, Hz), 7. 88 (1H, d, J=8, Tz), 8. 61 (1H, d), 8. 71 (2H, a), 4. 16-4. 7. 16 (1H, d), 3. 87 (3H, a), 4. 16-4. 7. 16 (1H, d), 2. 02-1, 98 (1H, d), 1. 70-1, 60 (1H, d), 1. 5.	1H NAR (&) Dym 3004Hz, DASO-d6

Example No.	WS.	Purity >90	
	520 (H+1)	>90% (NMR)	Q ^T
179		1R)	3
IH NAR(8)	-1 15(7H m)	~	

Example No. 178 IH NHR(8) ppm 300MHz, DMSO-d6

8 ·

MS 582 (A+)	Purity >90% (NMR)	٥		91	Example No. 182
		H, s), 4. 26(1H, m), 2. 35-2. 1 5(2H, m), 2. 00-1. 75(4H, m), 1. 74-1. 55(1H, m), 1. 50-1. 1 5(3H, m)	(H, m), 8. 21 (1H, s), 7. 95 (1H, d, J= , d, J=8. 4Hz), 7. 86 (1H, d, J= 7. 8Hz), 7. 68-7. 56 (7H, m), 7 .14 (2H, d, J=8. 7Hz), 5. 21 (1	300MHz, DMSO-d8 8. 55(1H, d, J=2. 1Hz), 8. 32(IH NKR(8) ppm

	MS . 547 (M+1)
	Purity > 90% (NMR)
300kHz, pMSO-d6 8. 19(1H, s), 7. 95(1H, d, J=8 7. 7k2), 7. 86(1H, d, J=9, 7tz) 7. 65(4H, d, J=7, 4Hz), 7. 47 (2H, d, J=9, 7tz), 7. 44-7, 27 (6H, m), 6. 99(2H, d, J=8, 7tz) 1, 4. 20(1H, m), 2. 34-2, 12(2 11, m), 1. 98-1, 76(4H, m), 1. 6 4(1H, m), 1. 46-1, 13(3H, m).	
IH NMR(8) ppm	Example No. 181

EP 1 162 196 A1

EP 1 162 198 A1

Purity >90% (NMR) 605 (24+1)

855 054 (M+1)	
Example No. 186	ih Mar(8) ppm

MC Esa(M+1)	Purity >90% (NMR)		Example No. 185	
		000Hb., DMSO-46 8. 30 (1H, m), 8. 24 (1H, d, J=9, OHz.) 0Hz.) 8. 03 (1H, d, J=9, OHz.) 7. 79-7. 10 (9H, m), 5. 20-5. 07 (2H, m), 4. 43-4. 04 (4H, m) 3. 50-3. 36 (2H, m), 2. 40-1.	5 IH NAR(6) ppm	

B

Purity

>90% (NMR)

184

IH NAR(6) ppm

Example No.

MS 555 (M+1)	Purity >90% (NMR)		Example No. 188	
		30/MHz, DMSD-06 12.77 (1H, bra), 8, 21 (1H, d, J=1, 4Hz), 7. 92 (1H, d, J=0, 7 Hz), 7. 88 (1H, dd, J=0, 7, 1, 4 Hz), 7. 88 (1H, dd, J=0, 7Hz), 7 57-7. 27 (7H, m), 7. 11 (2H, d J=0, 7Hz), 5. 07 (2H, q), 4. 2 6 (1H, m), 2. 38-2. 16 (2H, m), 1. 98-1. 76 (4H, m), 1. 64 (1H, m), 1. 44 (1H, m)	IH NAR(8) ppm	

8

8

99-1, 57 (5H, m), 1.	MS 520 (N+1)
8), 5. 20 (2H, 6), 4. 31 (1H, br 1. J=12. 2Hz), 2. 35-2. 19 (2H	Purity >90% (NMR)
) 6. 90 (1H, d, j=8, 3Hz), 6. 8 3(1H, 9), 6. 74(1H, d, j=8, 0H	(
2(4H, A' B' q, J=8. 6Hz), 7. 53 (2H, d, J=7. 8Hz), 7. 37(1H, t	
d7. 86 (2H, ABq, J=8. 2Hz), 7.	
300MHz, DMSO-d6 12. 76 (1H, a), 8. 67 (1H, d, J=	* 0
IH NUR(8) ppm	Example No. 187

õ

Example No.

190

1H NMR(6) ppm

300MHz, DMSO-d6

8. 36-7, 90 (5H, m), 7. 74 (2H, d, 1-8, 6Hz), 7. 60-7. 40 (5H, m), 7. 25 (2H, d, 1-8, 7Hz), 5. 14 (2H, s), 4. 45-4. 28 (1H, m), 2. 40-2. 15 (4H, m), 1. 75-1. 55 (1H, m), 1. 65-1. 20 (3H, m)

Table 53

EP 1 162 196 A1

EP 1 162 196 A1

		·							
SW	Purity		Example N	MS	Purity	\$	on etdurexa	NS.	Antity
554 (H+1)	>90% (NMR)		No. 192	514 (H+1)	>90% (NMR)		0. 191	580 (M+1)	>90% (NMR)
	•	. 4Hz), 7, 86 (1H, d, J=8, 7Hz) 7, 61 (2H, d, J=6, 7Hz), 7, 26 7, 01 (6H, m), 4, 84 (2H, s), 4 .31 (1H, m), 3, 36 (4H, m), 2, 2 9 (2H, m), 2, 00-1, 75 (4H, m), 1, 75-1, 15 (10H, m)	1H NAR(6) ppm 300MHz, DMSO-d6 8. 22(1H, s), 7. 94(1H, d, J=8			11, 8). 7 , 7. 85(1) (24, d. J. (64, b.). (64, b.). 1, 2. 29 44, b.). 1. 50–1. 15	1H NAR(8) ppm		

Table 54

SW	Purity >		Example No.
580 (H+1)	>90% (NMR)		. 195
		300MH, DMSO-d6 8. 25 (1H, s), 7. 65 (2H, d, m), 7. 77 (1H, s), 7. 65 (2H, d, J-8, 4Hs), 7. 59-7. 51 (3H, m) 7. 43 (2H, d, J-8, 4Hs), 7. 17 (2H, d, J-8, 7Hs), 5. 10 (2H, s)), 4. 30 (1H, m), 2. 40-2. 16 (2H, m), 1. 7 5-1. 55 (1H, m), 1. 55-1. 10 (3H, m), 1.	1H NMR(8) ppm

N.S.	Purity > 9 (Example No.
524 (H+1)	>90% (NMR)		194
		300Hi, biso-d6 12.80(1, bes), 8.23(1H, s) 17.97(1H, d, J=8.5Hz), 7.87 (1H, d, J=8.5Hz), 7.70-7.17 (9H, m), 4.60-4.13(4H, m), 3.72-3.40(2H, m), 2.40-1.15 (14H, m)	IH NUR(8) ppm

	S 560 (M+1)	S.
	Purity >90% (NMR)	
3.00M/L, DMSO-d6 13.00(1H, bar), 8. 29(1H, d, 1-6. 8) 1-1.4Rz), 8. 15(1H, d, J-6. 8) He), 7. 97(1H, dd, J-1.4Rz, 8) He), 7. 89(2H, d, J-6. 8Rz), 7. 89(2H, d, J-6. 8Rz), 7. 80-7. 60(6H, m) 7. 26(2H, d) 1-8. 8Hz), 4. 47-3. 90 (4H, m), 3. 20-3. 10 (2H, m), 2. 41- 1. 22(14H, m)	04.5	
IH NMR(8) ppm	Example No. 193	T
		ı

EP 1 162 186 A1

EP 1 162 196 A1

					:		
MS 604 (M+1)	Purity >80% (NMR)		Example No. 198	MS 554 (H+1)	Purity >90% (NMR)		Example No. 197
19 (SH, D), 1. 40-1.	.₹,	300Hz, DMSO-d6 12.75(1H, 8), 8.23(1H, d, Je 4.4hz), 7.95and7.86(2H, AB d, Je, 6Hz), 7.69and7.19(4 H, A' 8 a, Je, 6hz), 7.36(1H, t, Je7.8Hz), 6.82(1H, d, Je 9.3Hz), 6.73(1H, e), 6.71(1 H, d, Je7.2Hz), 4.30(1H, bt- Je12.2Hz), 3.89(2H, d, Je3.21, 7Hz	1H NAR(8) ppm		:	300MHz, DMSO-d6 8. 23(1H, g), 7, 95(1H, d, J=8, 74a), 7. 86(1H, d, J=8, 74a) 7. 86(1H, d, J=8, 74a) 7. 59and7. 18 (4H, ABq, J=8, 77a), 7. 35(1H, t, J=8, 44z), 8. 80-6, 70(3H, m), 4. 82(2H, g), 4. 31 (1H, m), 3. 40 (4H, m), 2. 29(2H, m), 2. 00-1, 75(4H, m), 1. 70-1. 15(10H, m)	1H NMR(8) ppm

	_			
MS.	Purity		Example No	
514 (H+1)	>90% (NMR)		•	Tab.
			196	Table 56
•	-	3000Hz, DASO-46 8.22(IH, 5), 7, 55(IH, d, J=8, 4Hz) -4Hz), 7, 86(IH, d, J=8, 4Hz) 7, 59and7, 18 (4H, ABq, J=8, 7Hz), 7, 34 (IH, t, J=8, 0Hz), 6, 80-6, 69 (JH, m), 4, 83 (2H, s), 4, 31 (IH, m), 2, 98 (3H, s), 2, 29 (2H, m), 2, 28 (4H, m), 1, 70-1, 55 (1H, m), 1, 50-1, 15 (3H, m)	1H NMR(6) ppm	

Ŕ

	MS 553 (N+1)
0-1. 21 (3H, B)	Purity > 90% (NMR)
J=8. 2H ₂), 5. 14 (2H, 8), 2. 4 0-2. 19 (2H, m), 2. 04-1. 78 (4 H, m), 1. 71-1, 60 (1H, m), 1. 5	
-8. Thz), 7. 47(4H, 6), 7. 38(1H, t, J=8. 2Hz), 7. 20(2H, d, J=8. 7Hz), 6. 90(1H, d, J=8. 2 H-) 8 83(1H, c) 8 74(1H, d)	
(DMSO-d6) 8:8.28(1H, s), 8 .06(1H, d, J=8.7Hz), 7.92(1 H d T=8.7Hz) 7.72(2H d T	:0
1H NAR(8) ppm	Example No. 201
	MS 553 (4+1)
) 99 (1H, m), 1. 50-1. 20 (3H, m	Purity >90% (NMR)
),1.99-1.80(44.11),1.71-1	,

Example No.

204

1H NMR(6) ppm

300MHz, DMSO-d6 8. 39-8. 28(2H, m), 8. 08(1H, d, j=8. 8Hz), 7. 76(2H, d, j=8 .7Hz), 7. 29(2H, d, j=8. 7Hz), 7. 25-7. 13(2H, m), 6. 80-6. 60(3H, m), 4. 46-3. 98(4H, m) .3. 51-3. 42(1H, m), 3. 20-3. 04(1H, m), 2. 39-1. 20(14H, m)

Purity

>90% (NMR)

541 (M+1)

Example No.

203 203

300Miz, DuSO-d6 12. 74(1H, brs), 21 (1H, s) 12. 74(1H, brs), 21 (1H, s) 8. 88 (2H, d, J=9, 0Hz), 7, 93 (1H, d, J=3, 7Hz), 7, 85 (2h, d, J=8, 7Hz), 7, 58 (2H, d, J=8, 7Hz), 7, 13 (2H, d, J=9, 6Hz), 4, 50-4, 08 (4H, m) 3, 68-3, 30 (2H,

8 Purity Example No. >90% (NMR) 537 (M+1) 202 (DMSO-dB) & :12.81 (1H, brs), 8.24 (1H, s), 7.99 (1H, d, J) = 8.71 (2H, d, J) = 8.72 (2H, d, J) = 8.81 (2H, t, J) = 8.21 (2H, d, J) = 8.2 1H NAR(8) ppm IH NAR(8) ppm

EP 1 162 198 A1

EP 1 162 196 A1

124

Purity

>90% (NMR)

MS 558 (H+1)	Purity '>90% (NMR)		Example No.
	5-1. 15 (3H, a).	B. 6 E. 38. 29. 6	206 IH NMR(8) ppm
Ĺ		50200000000000000000000000000000000000	

¥

Purity >90% (NMR)

MS 513(H+1)

Example No.

210

1H NAR(&) ppm

	MS 653 (M+1)
	Purity >90% (NMR)
300MHz, DISO-d6 9. 59 (1H, br s), 8. 23 (1H, s), 9. 64 (1H, d, J=8. 4Hz), 7. 90 (1H, d, J=8. 4Hz), 7. 62 (2H, d, J=8. 7Hz), 7. 39 (2H, 2H, d, J= 8. 7Hz), 7. 18 (2H, d, J=6. 7Hz), 6. 63 (2H, d, J=8. 7Hz), 3. 95 -3. 37 (4H, m), 3. 51-3. 40 (1H -a), 3. 17-3. 92 (1H. m), 2. 39 -1. 18 (17H, m)	- de de de
205 IH NMR(6) ppm	Example No.

Table 59

EP 1 162 196 A1

X.	10	·	m
S	Purity	= .	Example No.
			e z
6	> 9 0	OT	ļ.
582 OI+1)	% ()	Q	
1)	>90% (NMR)		
	٥	S	208
	لِـا	COTO: Timbi	
	표.	0Hz). 0Hz). 19-7. 7(18. 1Hz 1-2. 15(1-2. 15(1. 15).	IH NAR(6) ppm 300MHz, DMSO-d6
		77-8.00 77.40(2) 7.40(2) (2H, d, J 1.8), 2. 1.9), 1.	26 PB
		25 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	È
		15.02.4 1.00.2.4 15.00.2.4	- เ

Example No.

209

TH NAK (6) Days

EP 1 162 196 A1

128

ਲੋ

Purity

>90% (NMR) 587 (H+1)

å

MS 575 (M+1)	Purity >90% (NMR)	+0-40-40+	Example No. 212
H, m), 1. 28 (9H, 8)	. <u>g</u> :⊢	300Miz. 0MSO-d6 8. 22(1H, s), 7. 93and7. 87(2) H. MBq. J=6, 6Hz), 7. 88and7, 17(4H, Å' B' q, J=6, 7Hz), 7. 4 3-7. 33(6H, m), 6. 87(1H, d, J -8. 1Hz), 7. 18(2H, d, J=8, 4H s), 6. 91(1H, d, J=6, 0Hz), 6. 81(1H, s), 6. 43(1H, b, J=6, 0Hz), 6. 181(1H, s), 4. 36(1H, b, J=6, 0Hz), 6.	1H NAR(8) ppm

MS 540 (M+1)	Purity >90% (NMR)	4090	Example No.
		ē (211
(KH =)	4. 8Hz), 2. 38-2. 20 (2H, m), 2	300MB, DMSO-68 8. 29(11), 8, 15and7. 47(2 H, ABq, J=9, 01b.), 7, 77and7. 24(41), ABq, J=8, 91b.), 7, 39(111, t., J=7, 241b.), 6, 84(11), 4, J=9, 31b.), 6, 76(11), 9, 6, 76 (111, d., J=9, 51b.), 4, 36(11), 9, 10, 10, 10, 10, 10, 10, 10, 10, 10, 10	IH NAR(6) ppm

EP 1 162 198 A1

EP 1 162 198 A1

Purity >90% (NMR) MS 519(M+1)		Example No. 216	Burity > 90% (NMR) MS 623(M+1)		Example No. 215	Purity > 9 0% (NMR) MS 490 (M+1)		Example No. 214	Table (
), 1, 66 (IK, m), 1, 50-1, 21 (3 H, m).	300HFs, MSO-do 12.77(H, E), 8. 23(LH, d, J= 1.4Rz), 7. 95(1H, d, J=8. 6Hs) 7. 7. 95(1H, dd, J=8. 6, 1.4Hs) 7. 7. 95(1H, dd, J=8. 61, 1.7. 61, 1.7. 62Hs), 7. 68, 7. 7. 67, 7. 68, 67, 7. 68, 7. 7. 60(H, a), 7. 261, 7. 10(1H, a), 7. 33(2H, d, J=8. 8tz), 7. 34(H, a), 7. 35(2H, d, J=8. 8tz), 4. 31 (1H, a), 3. 80(2H, a), 2. 48-2 (1H, a), 3. 80(2H, a), 2. 48-2	1H NAR(8) ppm	, n), 1.49-1, 18 (3H, n).	12.75(11, tns) 8.23(11, s) 12.75(11, tns) 8.23(11, s) 17.95(11, d, J=8, Ths), 7.85 (11, d, J=8, Ths), 7.75(21, d) J=8, 4hs), 7.7(21, d, J=8, d) 4.7, 7.63-7.39(21, m), 7.5 (21, d, J=8, 4hs), 7.24(21, d) 4. J=8, 4hs), 7.18(11, s), 4.31(11, m), 4.33(11, m), 4.33(11, m), 4.33(11, m), 4.39-2.20(21, m) 2.00-1.76(41, m), 1.65(11	2 00 P), 4, 31 (1H, a), 2, 38-2, 19 (2 H, a), 2, 00-1, 78 (4H, a), 1, 6 6 (1H, a), 1, 48-1, 22 (3H, a).	i, d., J=2 1Hz), 8. 6 j=4. 8, 1.5Hz), 8. 12 (1H, i, 6Hz), 8. 12 (1H, 2. 1Hz), 7. 93 (1H, 2. 1Hz), 7. 90 (1H, d., J= j), 7. 70 (1H, d., J= j-7. 7. 54 (3H, m.), 7. j=8. 1, 4. 8Hz), 7. 21 (1H	1H NUR(8) ppm 300MHz, DMSO-d6	62

ŧ

ផ្ល

5 Purity

>90% (NMR)

544 (M+1)

Ġ

6

Example No.

219

IH NER(8) ppm

ĸ

Purity"

>90% (NMR)

¥

558 (H+1)

S

MS	Purity		Example No.
554 (M+1)	>90% (NMR)		0. 221
	9(4H, m), 1. 70-1. 59(1H, m), 1. 44-1. 20(3H, m)	8.23 (14, s), 7.96and7.86(2 H, ABq, JES, 6Hz), 7.69and7. 18 (4H, A' B' q, JeS, 7Hz), 7.3 7 (1H, t, JeS, 2Hz), 6.87 (1H, s), 6. 75 (1H, d, JeS, 0Hz), 5.24 (2H), 4.32 (1H, brt, Je12.2Hz), 4.32 (1H, s), 4.32 (1H, s), 2.30 H, m), 2,30 (3H, s), 2.00-1, 7	1H NHR(6) ppm

8

25

Example No.

218

IH NAR(&) ppm

8

300MHz, DMSO-d6 12. 9(1H, brs), R. 25(1H, s), 8. 04 (1H, d, 1-8, 7Hz), 7, 91(1H, d, 1-8, 6Hz), 7, 72(2H, d, 1-8, 5Hz), 7, 77(2H, d, 1-8, 7Hz), 7, 56 (2H, d, 1-8, 5Hz), 7, 62(2H, d, 1-8, 5Hz), 7, 62(2H, d, 1-8, 5Hz), 7, 13H, s), 2, 40-2, 15 (2H, m), 2, 71 (3H, s), 2, 40-2, 15 (2H, m), 2, 165 (1H, m), 1, 55-1, 15 (3H, m).

8

5

Purity

>90% (NMR)

602 (H+1)

ä

õ

(DMSO-d6) 6:12.80 (1H, brs), 8.23 (1H, s), 8.04 (1H, d; J), 8.05 (2H, s), 8.05 (2H, d; J=8, 6H, s), 7.96 (3H, d, J=8, 6H, s), 7.86 (1H, d, J=8, 7Hz), 7.763 (2H, d, J=8, 6Hz), 5.50 (2H, s), 4.36-4.21 (1H, m), 3.27 (3H, s), 2.47 (3H, s), 2.47 (3H, s), 2.47 (3H, s), 2.47 (3H, s), 1.77 (4H, m), 1.7

Example No.

217

IH NMR(8) ppm

MS 540 (M+1)	Purity >90% (NMR)		Example No. 220
. 8)	-1, 59 (1H, m), 1, 58-1, 20 (3H	12.76(1H, s), 8.23(1H, s), 7 12.76(1H, s), 8.23(1H, s), 7 96and7, 119(4H, A' B' q, 1-8, 9H, 2), 7.69and7, 119(4H, s), 1(1H, d), -7, 155(1H, s), 5, 1(1H, d), -7, 154(1H, d, 1-7, 154x), 6.85(1H, s), 6.7, 144(1H, d, 1-7, 154x), 5, 112, 124x) 12.65(3H, s), 2.41-2.20(2H, s), 2.65(3H, s), 2.41-2.20(2H, s)	1H NMR(6) ppm

EP 1 162 196 A1

Table 63

EP 1 162 196 A1

55

Purity

>90% (NMR)

580 (H+1)

23		50	43	40
MS.	Purity		ģ	Example No.
567 (M+1)	>90% (NMR)	ر <u>ڳ</u>	>-{} >-{}	
				228
		. 70 (2H, d, J=8. 7Hz), 4. 35-3 .97 (4H, m), 3. 62-3. 11 (2H, m), 2. 96 (6H, s), 2. 39-1. 12 (1 4H, m)	IH, d. Jeg Jeg, gas) Hz), 7, 32	1H NUR(6) ppm 300MHz, DUSO-d6 12 RO(1H hrs) 8 22(1H s)

NS.	Purity	-	ō Ex	1
564 (H+1)	:y > 90% (NMR)		Example No.	
m), 1. 49-1. 19(3H, m)	1, 75 (4H, m), 1, 73-1, 57 (1H,	A B q. j=8. 6kb, 7. 57 (14, 6), 7. 47 (14, d. j=6. 0ks), 7. 57 (14, d. j=6. 0ks), 7. 40 (24, t. j=8. 24s), 6. 91 (14, d. j=8.), 6. 95 (14, s.), 6. 15 (14, d. j=7. 9ks), 6. 26 (14, d. j=7. 9ks), 7. 27 (14, d. j=7. 9ks), 7. 28 (14, d. j=7. 9ks), 7.	227 1H NAR (6) ppm 300MHz, DMSO-d6 8, 43 (IH, d, J=6, 0Hz), 8, 23 (1H, 8) 7, 7, 96and 7, 18 (4H	

ㅂ

Purity

>90% (NMR)

(MSO-d6) 8:8, 22(1H, s), 8
.07(1H, d, J=8, 4Hz), 7:92(1
.H, d, J=8, 4Hz), 7:64(2H, d, J
.e., 716(2H, d, J=8, 4Hz), 7:10(2H, d, J=8, 4Hz), 7
.14(2H, d, J=8, 4Hz), 7
.14(2H, d, J=8, 32(1H, b), 3. 862
.(1H, hrd, J=12. 3Hz), 3. 65-3
.47(2H, m), 3. 10 (brdd, J=8, 4, 12. 3Hz), 2. 40-2. 20(2H, m), 2. 99-1, 76(6H, m), 1. 71-1
.16(6H, m)

544 (H+1)

5

Purity

>90% (NMR)

544 (X+1)

Example No.

224

IH NAR(6) ppm

S

Example No.

225

18 NMR(8) ppm

	SIL	Purity	, j	Example No.
N.	544 ()4+1)	>90% (NMR)	04	No.
2027		īŖ)	\$	226
an am (e)			300Mh. DMSO-6 8. 33ands. 08 (2H, ABq., J=8.7 Hz), 8. 31 (1H, m), 7. 66and7. 26 (4H, A B'q., J=9. 2Hz), 7. 4 2and7. 39 (4H, A' B'q., J=8.7H z), 4. 57 (2H, s), 4. 60 (1H, br t, J=12. 2Hz), 3. 85–3. 62 (3H m), 3. 28–3. 16 (2H, m), 2. 42 -2. 23 (2H, m), 2. 14–1. 81 (6H m), 1. 72–1. 25 (6H, m)	1H NAR(8) ppm

EP 1 162 196 A1

Table 65

223

1H NMR(8) ppm

300HHz, DMSD-d6 10. 96(1H, brs), 21 (1H, d, 10. 94(1H, brs), 21 (1H, d, 7-9.4 Hz), 7. 93(1H, d, J-6, 7, 1, 4 Hz), 7. 84(1H, dd, J-6, 7, 1, 1, 4 Hz), 7. 76-7. 40(7H, m), 7. 18 (2H, d, J-6, 0Hz), 4. 24-4, 16 (2H, m), 2. 40-1, 12 (18H, m)

Example No.

132

EP 1 162 196 A1

Example No. 231 IH NAR(6) ppm 300Mtz DMSO-d6 12.78(1H, brs), 8.23(1H, brs), 8.23(1H, brs), 8.23(1H, brs), 7.87(1H, d.), 7.87(1H, d.), 7.87(1H, d.), 7.87(1H, d.), 7.87(1H, d.), 7.87(1H, d.), 8.41z, 7.75(2H, d.), 8.41z, 7.75(2H, d.), 8.41z, 7.75(2H, d.), 8.41z, 7.75(2H, d.), 8.41z, 7.24(2H, d.), 8.22(2H, d.), 8			
11 NMR (6) ppm 300Mbz DMSO-d6 12. 78 (1H, brs), 8. 23 (1) 19-1. 5Hz), 7. 96 (1H, d. J. 19-1. 5Hz), 7. 76 (1H, d. J. 19-1. 5Hz), 7. 76 (2H, d. J. 19-1. 52 (2H, d. J. 19-1. 52 (3Hz), 7. 24 (2H, 19-1. 54 (2Hz), 7. 24 (2Hz)	Purity about 90% (NMR)	i di	
Dpm SO-d6 SO-d			231
8. 7 1. 5 2. 7 2. 7 2. 7 2. 7 2. 7 2. 7 2. 7 2. 7	(元), 1.49-1.17(3H, 元)	0MHz DMSO-d6 .78(1H, brs), 8, 1.5Hz), 7.96(1H, 1.787(1H, dd, J) 7.75(2H, d, J= 3, 7.75(2H, d, J= 4, J=8, 4Hz), 7, 4Hz), 6, 4Hz), 7, 4Hz), 6, 4Hz), 7, 8), 2, 97(6H, br	1H NAR(6) ppm

77 (1H, d d, J=9. (7H2), 77. 64 (7H2), 77. 60 (7H2), 15 (2H	1. 55-1. 20 (3H, m).		Example No. 230 Purity about 90% (NMR)	(, 6) pi , Diso- H, s), 8 , 05(1) (2H, d, J J-8, 11 J-8, 11 J-8, 21 J, 22, 40 J, 22, 40 J, 176 J, 17
--	----------------------	--	---	---

(1440809 SW	Purity > 90% (NMR)	0		°	Example No. 229
		9Hz), 4, 30-4, 20 (1H, m), 2, 3 9Hz), 4, 30-4, 20 (1H, m), 2, 3 8-2, 18 (2H, m), 1, 98-1, 18 (8 H, m), 1, 35 (3H, t, J=6, 9Hz)	92(1H, d, J=8, 1Hz), 7, 84(1H , d, J=9, 9Hz), 7, 62-7, 50(7H , m), 7, 12(2H, d, J=8, 7Hz), 5	300MHz, DMSO-d6 8. 25(1H, s), 8. 20(1H, s), 8. 04(1H, dd, J=8. 1, 1. 8Hz), 7.	IH NAR(6) ppm

EP 1 162 196 A1

Example No. 232 IH NMR(300Miz. 12.8 (IH 17.56 (IH 18.4 (
232 9 0 % (NMR)	5	Purity	<u></u>	Example
232	FO 809	>90%(00	
	Ė	NMR)	ific	232
), 1. 50-1	11, d, J=6 , 7, 59 (21) -7, 50 (51) -7, 942) / 7 , 942) / 7 , 5, 11 (2) 3, 01 (31, 3, 2, 40-2	32 IH NMR(6) ppm 300MHz, DMSO-de 12.8(1H, brs), 8

Example No.

239

1H NAR(6) ppm

13. 20(1H, br.s), 8. 89(1H, s), 8. 32(1H, s), 8. 25 (1H, d, J=8, 812), 18. 94 (1H, d, J=8, 812), 18. 94 (1H, d, J=8, 612), 7. 79-7. 74 (4H, m), 7. 60(2H, d), 7. 52(2H, d), 7. 52(2H, d), 4. 38 (1H, m), 2. 72 (3H, s), 2. 50-2, 15 (2H, m), 1. 75-1, 15 (3H, m), 1. 95-1, 76 (3H, m), 1. 75-1, 15 (3H, m), 1. 55-1, 15 (3H, m)

EP 1 162 196 A1

Ē

Purity S

> 9 0% (NMR) 538(M+1-2HC1)

SW Purity

>90% (NMR) 553(M+1-HC1)

Example No.

234

1H NAR(8) ppm

DMSO-d6 8. 77 (1H, d, J=3, 6Hz), 8. 36-8. 26 (3H, a), 8. 08 (1H, d, J=8, 6Hz), 7. 79 (2H, d, J=8, 7Hz), 7. 78 (2H, d, J=6, 7Hz), 7. 58 (2H, d, J=6, 4Hz), 7. 55 (2H, g), 4. 38 (1H, a), 5. 26 (2H, g), 4. 38 (1H, a), 2. 10 (2H, a), 1. 95-1. 76 (2H, a), 1. 95-1. 76 (2H, a), 1. 95-1. 15 (3H, a), 1. 75-1.
g

5

员

Purity

>90% (NMR)

APCI-NS 525 (N+1)

8

Ġ

ŧ

ä

ĸ

Purity

>90% (NMR)

APCI-Ms 622(M+1)

S

Example

o.

240

1H NMR(8) ppm

300MHz, DMSO-d6 8, 90(H, s), 8, 25(IH, d, J=8, 3) 29(IH, s), 8, 25(IH, d, J=8, 3) Hz), 8, 05(IH, d, J=8, Hz), 7 96(IH, s), 7, 93(IH, d, J=8, 4) J=8, 8Hz), 7, 83(IH, d, J=8, 4) Hz), 7, 697, 59(ZH, m), 7, 54 (ZH, d, J=8, 8Hz), 4, 37(IH, b) rt), 2, 30(ZH, m), 2, 00(ZH, m)), 1, 88(ZH, m), 1, 67(IH, m), 1, 5-1, 2(3H, m)

ક

300MHz, DMSO-46 12. 79 (1H, brs), 8, 60 (2H, d, J-1, SHz), 8, 53 (1H, s), 8, 25 (1H, s), 8, 25 (1H, s), 7, 98 and 7, 85 (2H, MB d, J-9, 4Hz), 7, 78 (2H, d, J-9, 4Hz), 7, 74 (4H, d, J-6, 6Hz), 7, 44 (4H, d, J-6, 6Hz), 7, 44 (4H, d, J-6, 6Hz), 7, 44 (4H, d, J-6, 6Hz), 7, 43 (2H, m), 7, 43 (2H, m), 2, 03-1, 82 (4H, m), 1, 72-1, 61 (1H, m), 1, 42-1, 22 (3H, m)

ğ

ß

8

ä

õ

MS APCI-Ms 655 (M+1)	Purity > 90% (NMR)		Example No. 236	
		3000Hz, DMSO-d-6 8. 40-7. 40 (11H, m), 2. 95, 2. 81 (3H, each d, J=4. 7Hz), 2. 40-2. 20 (2H, m), 2. 10-1. 80 (4H, m), 1. 70- 1, 60 (1H, m), 1. 50-1. 20 (3H, m)	IH NUR(6) ppm	

WS APCI-Ms 838 (H+1)	Purity >90% (NMR)		Example No. 2	
		12. 74(1H, brs), 8. 67(1H, dd 12. 74(1H, brs), 8. 67(1H, dd . J=3, 1, 1. 6Hz), 8. 21(1H, d, . J=1, 6Hz), 7. 93(1H, dJ=8, 6H z), 7. 90–7. 80(2H, d), 7. 60– 7. 50(7H, d), 7. 09(2H, d, J=6 7. 7Hz), 5. 16(2H, s), 4, 25 (1H . m), 2. 40–2. 20 (2H, d), 2. 00 -1, 60 (5H, d), 1. 50–1. 20 (3H . m)	235 1H NMR(6) ppm	

EP 1 162 198 A1

EP 1 162 196 A1

Example No. APCI-Ms 521 (M+1) 239 IH NAR(6) ğ

8

5 Purity >90% (NMR)

Example No. 238 300MHz, PMSO-46 12.80 (IH, brs), 8, 54 (IH, s) 12.80 (IH, brs), 8, 64 (IH, s) 8, 25 (IH, s), 7, 98 and 7, 88 (2H, Abq, J-8, 6Hz), 7, 76 (2H, d, J-8, 6Hz), 7, 53-7, 31 (3H, m. 6, 61 (IH, s), 5, 46 (2H, s) 4, 32 (IH, brt), 2, 40-2, 20 (2H, m), 2, 02-1, 79 (4H, m), 1, 69-1, 69 (IH, m), 1, 49-1, 19 (3H, m) 1H NMR(6) ppm

ផ្ល

!		1005	1003	1002	1001	Ex. No.
						Formula
	385 (M+H)	357 (M+H)	398 (M+H)	454 (M+H)	364 (M+H)	SW

EP 1 162 196 A1

8	8	å .	8	ક ક	8 6	8	u	
	1012	**************************************	1010 m 1777		1008		Table 72 Ex. No. Formula	EP 1 162 196 A1
	366 (M+H)	395 (M+H)	390 (M+H)	310 (H+H)	416 (M+H)	557 (C\$H)	NS.	

1018	1017	1016	1015	1014	1013	Ex. No.	
						Formula	Table /3
340 (M+H)	414 (M+H)	402 (M+H)	350 (M+H)	382 (M+H)	374 (M+B)	SW	

EP 1 162 198 A1
Table 73

EP 1 162 196 A1

4

1029	1028	1027	. 1026	1025	1024	Ex. No.	
						Formula	Table 75
362 (M+H)	366 (M+H)	408 (M+H)	336 (M+H)	408 (M+H)	218 (M+H)	SW	

Table 76

EX. No. Formula MS

1030

1030

1031

1031

1032

1032

1033

1033

1034

1034

1035

1035

1036

1037

1038

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

1039

EP 1 162 196 A1

EP 1 162 196 A1

4

1041	1040	1039	1038	1037	1036	Ex. No.
,						Table 77 Formula
440 (M+H)	417 (M+H)	406 (M+H)	466 (M+H)	428 (M+H)	412 (M+H)	MS

EP 1 162 198 A1

EP 1 162 198 A1
Table 78

1047	1046	1045	1044	1043	1042	Ex. No.	
						Formula	Table 78
307 (M+H)	352 (M+H)	423 (M+H)	312 (M+H)	440 (M+H)	417 (M+H)	SW	

1052	1051	1050	1049	1048	Ex. No.	
					Formula	Table /9
518 (M+H)	442 (M+H)	326 (M+H)	398 (M+H)	374 (M+H)	SW	

EP 1 162 196 A1

1058	1057	1056	5501	1054	1053	Ex. No.	
OH DE LA CONTRACTION OF THE CONT	POP COM	14-V-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1				Formula	EP 1 182 188 A1
367 (M+H)	367 (M+X)	352 (M+H)	442 (M+H)	376 (M+H)	442 (M+H)	SW	

\$90t	1063	1062	1061	1060	1059	Ex. No.
		***************************************	14x			Formula
351 (M+H)	360 (M+H)	357 (M+H)	352 (M+H)	324 (M+H)	364 (M+H)	SW

Ex. No. Formula MS

1065

1066

1067

1068

1069

1069

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

1070

EP 1 182 196 A1

5 5 5 8 5 6 5

1076	1075	1074	1073	1072	1071	Ex. No.	
						Formula	Table 83
386 (M+H)	337 (M+H)	358 (M+H)	399 (M+H)	455 (M+H)	365 (M+H)	SW	

EP 1 162 198 A1

1082	1081	1080	1079	1078	1077	Ex. No.	
						Pormula	TABLE 84
367 (M+H)	396 (M+H)	391 (M+H)	311 (M+H)	417 (M+H)	358 (M+H)	SH.	

ź

1088	1087	1086	1085	1084 .	1083	Ex. No.
					100	Formula Formula
341 (M+H)	415 (M+H)	403 (M+H)	383 (M+H)	351 (M+H)	375 (M+H)	SW

EP 1 162 196 A1

EP 1 162 196 A1

ź

1099	1098	1097	1096	1095	1094	Ex. No.	
	\$ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\					Formula	Table 87
363 (M+H)	367 (M+H)	409 (M+H)	337 (M+H)	409 (M+H)	519 (M+H)	SW	

1105	1104	1103	1102	1101 .	1100	Ex. No.	
				HO COH		Formula	EP 1 162 196 A1 Table 88
413 (M+H)	413 (M+H)	467 (M+H)	308 (M+H)	339 (M+H)	474 (M+H)	SW	

ŕ

Ē

	1110	1109	1108	1107	1106	Ex. No.	
						Formula ,	Table 89
418 (M+H)	441 (M+H)			467 (M+H)	429 (M+H)	MS	

EP 1 162 186 A1

EP 1 182 196 A1 Table 90 Formula

(H+M) 80E

1122	1121	1120	0 11	8 111	Ex. No.		
					Formula	Table 91	EP 1 162 196 A1
353 (M+H)	443 (M+H)	377 (M+H)	443 (M+H)	519 (M+H)	MS		

1128	1127	1126	1125	1124	1123	Ex. No.	
					HO HO	Formula	Table 92
358 (M+H)	353 (M+H)	325 (M+H)	365 (M+H)	368 (H+H)	368 (M+H)	SW	

ź

1134	1133	1132	1131	1130	1129	Ex. No.
						Formula
365 (M+H)	368 (M+H)	367 (M+H)	352 (M+H)	352 (M+H)	361 (M+H)	SW

EP 1 162 196 A1

1140	1139	1138	1137	1136	1135	Ex. No.	
						Formula	EP 1 162 188 A1 Table 94
387 (M+H)	467 (M+H)	365 (M+H)	385 (M+H)	307 (M+H)	351 (M+H)	SW	

1146	1145	1144	1143	1142	1141	EX. No.
						Formula
385 (M+H)	484 (M+H)	363 (M+H)	323 (M+H)	364 (M+H)	322 (M+H)	NS

EP 1 162 198 A1

1151	1150	1149	1148	1147	Ex. No.	
458 (M+H)	458 (M+H)	(H+M) 805	100 (M+H)	427 (M+H)	Formula	EP 1 162 198 A1 Table 96

Ē

	1154	1153	1152	Ex. No.	
				Formula	Table 97
454 (N+H)	508 (M+H)	458 (M+H)	474 (M+H)	SW	

EP 1 162 188 A1

Table 98

Formula

Formula

MC ON

495 (M+H)

MC ON

495 (M+H)

MC ON

495 (M+H)

MC ON

496 (M+H)

EP 1 162 196 A1

8	E	8	5	õ	

1165	1164	. 1163	1162	1161	Ex. No.	İ
	10) Come				Formula	EP 1 162 196 A1
420 (M+H)	526 (M+H)	466 (M+H)	447 (M+H)	468 (M+H)	SK	

1171	1170	1169	1168	1167	1166	Ex. No.	4	
406 (N+H)	404 (M+H)	136 (H+H)	436 (M+H)	135 (M+H)	490 (M+H)	Formula	8 41	

1177	1176	1175	1174	1173	1172	Ex. No.	
						Formula	Table 101
406 (M+H)	523 (M+H)	420 (M+H)	406 (M+H)	420 (M+H)	392 (M+H)	SW	

EP 1 162 198 A1

EP 1 162 198 A1

5 5 8 8 8 3 8 5

90 90 71	11 85	1184	1183	1182	Ex. No.	
					Formula	Table 103
490 (M+H)	508 (M+H)	418 (M+H)	496 (N+H)	497 (M+H)	NS.	

EP 1 162 198 A1

EP 1 162 198 A1

1198	1197	1196	1195	1194	1193	Ex. No.
						Formula
509 (M+H)	456 (M+R)	497 (M+H)	516 (M+H)	504 (M+H)	512 (M+H)	SW

Table 106

Ext. No. Formula MS

1199

1200

1200

1201

1202

1203

1204

1204

1204

1206

1207 (M+H)

1208

1209

1201

1201

1201

1202

1203

1204

1206

1207 (M+H)

1208

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

1209

EP 1 162 198 A1

EP 1 162 198 A1

1210 Ho		1208	1207	1206	45 CO 24 CO	
332 (M+H)	363 (N+H)	385 (M+H)	341 (M+H)	325 (M+H)	454 (M+H)	

EP 1 162 196 A1

EP 1 162 196 A1

1222	1221	1220	1219 .	1218	1217	Ex. No.	
						Formula	Table 109
332 (M+H)	341 (M+H)	385 (N+H)	337 (M+H)	391 (M+H)	433 (M+H)	¥5	

EP 1 162 196 A1

426 (M+H) 321 (M+H) (H+M) 15E

176

178

EP 1 162 198 A1

EX. NO. FORMULA MS

1235 MO 498 (M+H)

1236 MO 4 498 (M+H)

1237 MO 502 (M+H)

1238 MO 4 498 (M+H)

1239 MO 4 498 (M+H)

1239 MO 4 498 (M+H)

1239 MO 4 498 (M+H)

EP 1 162 196 A1

178

EP 1 162 198 A1

1245	1244	1243	1242	1241	1240	£x. No.	
						Formula	Table 113
454 (M+H)	494 (M+H)	468 (M+H)	46D (M+H)	408 (M+H)	483 (M+H)	SW	

1250 Table 114
Formula 460 (M+H) 468 (M+H)

1254	1253	1252	1251	Ex. No.	
				Formula	Table 115

EP 1 162 196 A1

Ŕ

=

Table 117
Formula

505 (M+H)

EP 1 162 186 A1

1266 1266

EP 1 182 198 A1

Ē

6 6 8 8 6 8

1270	1269	1268	1267	Ex. No.	
				Formula	Table 119
498 (M+H)	468 (M+H)	454 (M+H)	494 (M+H)	SK	

Table 120

FORMULA

1271

1272

1272

1273

1274

1274

1274

FORMULA

FORM

EP 1 162 186 A1

EP 1 162 198 A1

88

Ē

1278	1277	1276	1275 Ho	Ex. No.
				Formula
516 (M+H)	456 (м+н)	427 (N+H)	519 (M+H)	MS

Table 122

Ex. No. Formula N3

1279

1279

1280

AS (N+H)

1281

AS (N+H)

1282

AS (N+H)

É

187

EP 1 162 198 A1

6 2 8 2 3 6 2

1288	1287	1 22 8 G	1 N 00 5	1284	Ex. No.	
					Formula	Table 123
508 (M+H)	508 (M+H)	420 (M+H)	406 (M+H)	482 (M+H)	MS	

EP 1 162 196 A1

Ex. No. Formula

1289

1289

1290

1291

1292

1292

EP 1 162 196 A1

ź

1297	1296	567 t	1294	1293	FX. No.	
					Formula	Table 125
470 (3+H)	908 (N+H)	477 (H+H)	496 (M+H)	490 (M+H)	SW	

EP 1 162 196 A1

2

					_
1306	1304	1303	1302	Ex. No.	
8				Formula	Table 127
440 (M+H)	504 (M+H)	399 (N+H)	51.3 (H+H)	NS	

1311		1309	1308	1307	Ex. No.	
					Formula	EP 1 162 186 A1
522 (M+H)	532 (M+H)	218 (M+H)	508 (M+H)	494 (M+H)	SW	

Ŕ

1 .

1315	1314	13 13 14	1312	Ex. No.	
				Formula	EP 1 162 196 A1
488 (M+H)	517 (M+H)	484 (N+H)	546 (M+H)	MS	

3 22	1321	1320	1319	1318	1317	Ex. No.	
						Formula	Table 130
522 (M+H)	522 (M+H)	510 (M+B)	504 (N+H)	423 (N+H)	413 (M+H)	SM	

2

ĝ

1327	1326	1325	132 4	ր ա Խ Ն	Ex. No.
					Formula
496 (M+H)	491 (M+H)	502 (M+H)	449 (M+H)	4 84 (X+H)	MS

) i i i o o

EP 1 162 186 A1

198

1338	1337	1336	1335	1334	1333	
			"TOHO!"		Formule	Table 133
498 (M+H)	484 (M+H)	484 (M+H)	547 (N+H)	536 (H+H)	MS 164	

EP 1 162 196 A1

Table 134
Formula 502 (M+H) (H+M) 88P 513 (M+H) 514 (M+H) 498 (M+H) (H+M) 825

1350	1349	1348	1347	1346	1345	Ex. No.	
						Formula	EP 1 152 186 A1 Table 135
546 (M+H)	522 (M+H)	480 (M+H)	499 (H+H)	502 (M+H)	488 (M+H)	SW	

1356	1355	1354	i u u	1352	1351	Ex. No.	
						Formula	Table 136
566 (M+H)	4B0 (M+H)	532 (M+H)	609 (H+H)	484 (M+X)	482 (M+H)	SW	

EP 1 162 196 A1

1368	1367	1366	1365 13	1364	1363	Ex. No.	
						Formula	Table 138
497 (M+H)	524 (M+H)	481 (M+H)	488 (M+H)	494 (M+H)	512 (M+H)	SW	•

EP 1 182 196 A1

1374	1373	1372	1371	1370 .	1369	Ex. No.
5						Table 139 Formula
458 (M+H)	494 (M+H)	469 (M+H)	470 (M+H)	469 (M+H)	472 (M+H)	MS

EP 1 162 196 A1

1380	1379	1378	1377	1376	1375	Ex. No.	
						Formula	Table 140
\$10 (M+H)	496 (M+H)	526 (M+H)	542 (M+H)	554 (M+H)	612 (M+H)	SW	

EP 1 162 196 A1

	1384	7 08 04 14	1382	1381	Ex. No.	
					Formula	Table 141
539 (M+H)	523 (M+H)	558 (M+H)	525 (M+H)	540 (M+H)	SW	

EP 1 182 198 A1

1396	بر ب پ پ	1394	1393	1392	1391	Ex. No.
						Formula
506 (M+H)	491 (M+H)	507 (M+K)	525 (M+H)	527 (M+H)	540 (M+H)	SM

522 (M+H)

(R+H) 8ES

(H+M) 0EG

EP 1 162 196 A1

Table 144
Formula

522 (M+H)

EP 1 162 196 A1

1408	1407	1406	1405	1404	1403	Ex. No.	
					**************************************	Formula	Table 145
547 (M+H)	469 (M+H)	455 (M+H)	472 (M+H)	475 (M+H)	534 (M+H)	SW	

Table 146

Ex. No. Formula MS

1409

1410

1411

1411

1411

1411

1411

1411

1414

1414

1414

1414

1416

Formula MS

252 (M+H)

504 (M+H)

504 (M+H)

522 (M+H)

468 (M+H)

EP 1 162 198 A1

EP 1 162 196 A1

1420	1419	1418	1417	1416	1415	Ex. No.
						Table 147 Formula
522 (M+H)	455 (M+H)	455 (M+H) -	502 (M+H)	488 (M+H)	502 (M+H)	MS

EP 1 162 196 A1

1425	1424	1423	1422	1421	Ex. No.	
					Formula	EP 1 162 195 A1
458 (M+H)	494 (M+H)	S10 (M+H)	536 (M+H)	469 (м+н)	SW	

1430	1429	1428 2	1427	 Ex. No.	
				Formula	Table 149
511 (M+H)	441 (M+H)	480 (M+H)	526 (M+H)	MS MS	

EP 1 162 196 A1

EP 1 162 198 A1

1442	1441	1440	1439	1438	1437	Ex. No.	
						Formula	EP 1 162 186 A1 Table 151
474 (M+H)	508 (M+H)	490 (M+H)	474 (M+H)	508 (M+H)	524 (M+H)	SW	

1446	1445	4 4 1	1443	1
10 (M+H)	4,c du, (M+H)	600 (M+H)	Formula MS 516 (M+H)	2 88 A1

Ŋ

1453	1452	1451	1450	1009		Ex. No.
						Formula
508 (H+H)	441 (M+H)	474 (M+H)	490 (H+H)	440 (M+H)	(37.2)	SW

EP 1 162 196 A1

1465	1464	1463	1462	1461	1460	Ex. No.
						Table 155 Formula
502 (M+H)	460 (M+H)	476 (M+H)	427 (M+H)	S16 (M+H)	486 (M+H)	SW

EP 1 162 196 A1

1470	1469	1467	Ex. No.
HO 512 (M+H)	2598 (M+H)	230 (H+H) 230 (H+H)	Table 156 Formula MS 586 (M+H)

1477	1476	1475	1474	1473	1472	Ex. No.
						Table 157 Formula
435 (M+H)	508 (M+H)	441 (M+H)	474 (M+H)	490 (M+H)	440 (M+H)	MS

EP 1 162 198 A1

14 B2	1481	1480	1479	1478	Ex. No.	
					Formula	Table 158
482 (M+H)	426 (M+H) -	516 (M+H)	496 (M+H)	522 (M+H)	NS	

EP 1 162 198 A1

1486	148 85	14 84 4	1483	Ex. No.	•
				Formula	Table 159
476 (M+H)	427 (M+H)	516 (M+H)	486 (M+H)	SW	

EP 1 162 198 A1

226

1494	1493	1492	1491	Ex. No.	
				Formula	Table 161
544 (M+H)	512 (M+H)	598 (M+H)	330 (8743)	MS	

EP 1 162 196 A1

6

1503	155 02	1501		Ex. No.
				Formula
656 (M+H)	600 (M+H)	630 (M+H)	606 (M+H)	NS.

EP 1 162 198 A1

1507		1506	1505	1504	Ex. No.	
	-layer Cy.	\$	140 Ch.	, -cy,	Formula	Table 164
580 (M+H)		656 (M+H)	(H+H) 009	630 (M+H)	SW	

EP 1 162 196 A1

1512	1511	1510	1509	1508	Ex. No.	
					Formula	Table 165
546 (M+H) ·	550 (M+H)	580 (M+H)	606 (M+H)	550 (M+H)	SW	

1521	1520	1519		Ex. No.	
				Formula	Table 167
606 (M+H)	628 (M+H)	572 (M+H)	602 (M+H)	SW	

1525

(H+M) 245

(H+M) 209

Table 168 Formula

SW

EP 1 162 196 A1

234

Ex. No. Table 169 Formula 628 (M+H) (H+M) P19

EP 1 162 198 A1

Ex. No. 1533 Table 170 Formula 584 (M+H) (H+H) P19 (H+W) 8 T9 (H+M) 885

235

(H+M) 909

SE

EP 1 182 196 A1

1537	1536	1535	Ex. No.	
			Formula	Table 171
627 (M+H)	627 (M+H)	640 (M+H)	SW	

Table 172

Ex. No. Formula MS

1538

1538

No. Formula MS

560 (M+H)

1539

No. Gar (M+H)

1540

1541

1541

1541

1541

1541

1541

ä

23

Table 174
Formula

EP 1 162 196 A1

246

) U). 55 2	<u>រ</u> ភ ភ	1550 .	Ex. No.
				Table 175 Formula
34 34 8	532 (M+H)	560 (M+H)	627 (M+H)	SK

EP 1 162 198 A1

1559	1558	Ex. No.	
		Formula	Table 177
570 (M+H)	584 (M+H)	MS	

[0292] The evaluation of the HCV polymerase inhibitory activity of the compound of the present invention is explained in the following. This polymerase is an onzyme coded for by the non-structural protoin region called NSSB on the RNA gene of HCV (EMBO J., 15:12-22, 1988).

Experimental Example (i)

â

i) Preparation of enzyme (HCV polymerase)

(0293) Using, as a template, a cDNA clone corresponding to the full length RNA gene of HCV BX strain obtained from the blood of a patient with hepatitie C, a region ancoding NSSB (661 amino acids; J Virol 1991 Mar, 55(3), 1105-13) was empilled by PCR. The objective gene was prepared by adding a 6 His tag (base pair encoding 0 continuous histidine (His)) to the 5' end thereof and transformed to Eacherfichis colf. The Eacherfichia colf capable of producing the objective protein was cultured. The obtained cells were suspended in a buffer solution containing a surfactant and crushed in a microfluidizer. The supermatant fives belanded by certificipation and applied to various column chromatography; for the control of the cells were supermatant and crushed in a microfluidizer. The supermatant fives belanded by certificipation and applied to various column chromatography; for the objective protein supermatant cells. give a standard enzyme product

ii) Synthesis of substrate RNA

8 [0294] Using a synthetic primer designed based on the sequence of HCV genomic 3' untranslated region, a DNA fregment (148 bp) containing polycl and 3'x sequence was entrely synthetized and cloned into pitamid p8uestrips ISK III+) (Stratagene). The cDNA encoding hull length NSSB, which was prepared in J above, was digested with SK III+) (Stratagene). The cDNA encoding hull length NSSB, which was prepared in Jabove, was digested with striction enzyme Kpni to give a cDNA fragment containing the nucleotide sequence of from the restriction enzyme

EP 1 182 198 A1

cleavage alto to the termination codon. This cDNA fragment was inserted into the upstream of 3' untranslated region of the DNA in pillusertip SK II(+) and ligated. The about 450 bp inserted DNA asequence was used as a template in the preparation of substrate RNA. This plasmatives cleaved immediately after the 3'X sequence, linearized and purified by phenol-chloroform treatment and athanol procipitation to give DNA.

[2393] RNA was synthesized (37°C, 3 hr) by run-off method using this purified DNA as a template, a promoter of pBlucecript SK II(+), MEGAscript RNA epithesia kii (Ambhon) and TZ RNA polymorase. Divised was added and the mixture was incubated for 1 hr. The template DNA was removed by decomposition to give a crude RNA product. This product was treated with phenol-chloroform and purified by effective properties on to give a crude RNA product. This product was treated with phenol-chloroform and purified by effective properties on the objective substrate RNA. (2398] This RNA was applied to formaldehyde denaturation agarose get electrophorosis to confirm the quality thereof and preserved at -60°C.

iii) Assay of enzyme (HCV polymerase) inhibitory activity

[0297] A test substance (compound of the present invention) and a reaction mixture (30 μ) having the following composition were reacted at 25°C for 69 min.

(180 composition were reacted at 25°C for 69 min.

(180 mixture to stop the reaction. The reaction mixture was left standing in loo for 15 min to insolubilize RNA. This RNA was trapped on a glass filler (Whatman GFC and the like) upon literation by surface. The filter was veshed with a solution containing 1% tichleroscotic acid and 0.1% softum pyrophosphate, washed with 90% eitenol and dried. A liquid schilliation occitail (Fackart) was acided and the redosactivity of RNA synthesized by the enzyme reaction was measured on a liquid caltiliation counter.

(1929) The HCV polymerase inhibitory activity (C₉₀) of the compound of the present invention was calculated from the values of radioactivity of the enzyme reaction with and without the test substance.

(2030) The results are shown in Tables 179 - 184.

25 Reaction mixture : HCV polymerase (5 μg/m) obtained in), substatic RNA (10 μg/m) obtained in i), ATP (50 μM), GTP (50 μM), UTP (2 μ/M), [5.6-λ-(μ.TP (48 C/mmol (Amarsham), 1.5 μCl) 20 mM Tds-HCl (pH 7.5), EDTA (1 mM), MgCl₂ (5 mM), NaCl (50 mM), DTT (1 mM), BSA (0.01%)

		8			8				Ĝ			ô				ដ			ક	
2	61	50	49	48	47	46	45	4	43	30	26	20	17	12	11	9	8	2	Ex. No.	
0.58	1.0	0.54	0.94	0.44	0.54	0.47	0.40	0.85	0.58	0.052	0.033	0.042	0.047	0.60	0.53	0.019	0.034	0.078	HCV polymerase inhibitory activity iCgo [நூ] Ex. No.	Теф
97	88	94	93	91	90	89	88	87	88	8.5	83	82	81	77	71	70	68	67	Ex. No.	Table 178
91.0	0.25	0.084	0.18	0.53	0.20	0.34	0.092	0.80	0.13	0.17	0.62	0.087	0.18	0.51	0.82	0.19	0.28	0.26	HCV polymerase inhibitory activity (C ₅₀ [µM]	

	;	.
230	229	Ex. No.
0.17	0.022	HCV polymerase inhibitory activity IC ₅₀ [μΜ]
259	267	Ex. No.
0.10	0.074	HCV polymerase inhibitory activity IC ₅₀ [µM]

Table 182

8	Ex. No. No.	HCV polymerase inhibitory activity IC ₂₀ E	Ex. No.	HCV polymerase inhibitory activity (C ₃₀ (µM)
1	183	0.03B	208	0.039
25	189	0.017	209	0.12
	190	0.020	210	0.31
	191	0.43	211	0.069
Ŗ	192	0.22	212	0.23
	193	0.13	213	0.10
	194	0.52	214	0.059
:	185	0.023	215	0.078
8	198	0.20	915	0.084
	197	0.11	217	0.058
	198	0.044	218	0.033
8	199	0.11	219	0.13
	200	0.10	220	0.073
	201	0.14	221	0.058
:	202	0.085	222	0.041
ŧ	203	0.083	223	0.21
	204	0.16	225	0.014
	205	0.077	227	0.045
8	208	80.0	228	0.18

Table 181

ě				7			•			
168	165	184	163	162	161	160	159	158	Ex. No.	
0.055	0.50	0.16	0.15	0.43	0.082	0.24	0.13	0.11	HCV polymerase inhibitory activity No. ICso [µM]	Table 180 (continued)
981	185	184	183	182	181	180	179	178	Ex.	ontinue
0.37	0.11	0.018	0.017	0.021	0.71	0.11	0.63	0.052	HCV polymerase inhibitory activity IC50 (µM)	Δ)

EP 1 162 196 A1

119 0.13 145 0.088

| Table 180 | Ex. No. HCV polymerase inhibitory activity No. IC₂₀ [µM] | Ex. HCV polymerase inhibitory activity No. IC₂₀ [µM] | Ex. HCV polymerase inhibitory activity No. IC₂₀ [µM] | Ex. HCV polymerase inhibitory activity IC₂₀ [µM] | 148 0.038 169 0.033 168 0.038 0.050 0.15 0.050 0.049 0.15 0.050 0.14 171 0.050 0.14 175 0.014 0.15 0.050 0.14 0.175 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.014 0.050 0.050 0.014 0.050

		Tabl	Table 179	
	Ex. No.	HCV polymerase inhibitory activity IC ₅₀ [μΜ]	Ex. No.	HCV polymerase inhibitory activity (C ₆₀ (µM)
0	88	0.53	120	0.16
	100	0.78	121	0.19
	101	0.14	122	0.51
13	103	0.17	123	0.10
•	104	0.073	124	0.091
	105	0.078	125	0.12
	108	0.40	128	0.14
٥	107	0.11	129	0.12
	108	0.21	130	0.18
	109	0.11	131	0.048
•	110	0.24	132	0.055
	111	0.14	133	0.12
	112	0.11	134	0.071
	113	0.071	139	0.26
۰	114	0.56	140	0.11
	115	0.17	141	0.43
	116	0.37	142	0.055
a	117	0.075	143	0.053
	118	0.14	144	0.19
	119	0.13	145	0.088

Table 178 (continued)

Ex. No. HCV polymerase inhibitory activity IC₂₀ (µM] Ex. No. HCV polymerase inhibitory activity IC₂₀ (µM] 55 0.38 98 0.30

	Ex. No.	HCV polymerase inhibitory activity IC30 [µM]	Ex. No.	HCV polymerase inhibitory activity (Ceo [µM]
Œ	283	0.014	298	0.011
	284	0.014	299	0.018
	285	0.012	300	0.045
	286	0.014	301	0.017
5	287	0.012	303	0.10
	288	0.013	304	0.017
	289	<0.01	305	0.01
G	290	0.012	306	0.013
	291	, 0.016	307	0.022
	292	0.015	308	0.023
	293	0.034	311	0.16
8	294	0.032	312	0.023
	285	0.045	313	0.025
	298	0.034	314	0.097
G	297	0.022	315	0.028

| Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | Ex. No. | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polymerase Inhibitory activity IC₂₀ [µM] | HCV polyme

EP 1 162 196 A1

| Ex. No. | HCV polymerase inhibitory activity IC₉₀ [µM] | Ex. No. | HCV polymerase inhibitory activity IC₉₀ [µM] | 316 | 0.022 | 502 | 0.024 | 317 | 0.032 | 503 | 0.198 | 318 | 0.012 | 601 | 0.32 | 0.32 | 319 | 0.030 | 701 | 0.052

(1+H) 919 SW	Purity > 90% (NMR)	3000	Example No.	•
	R)	1	250	
•		300KH, MISO-66 8. 26 (1H, d, J=1. 5Hz), 8. 16- 8. 08 (2H, m), 7. 99-7. 88 (2H, m), 7. 68 (2H, d, J=8. 6Hz), 7. 60-7. 48 (6H, m), 7. 19 (2H, d, J=8. 6Hz), 5. 17 (2H, g), 4. 31 (1H, m), 2. 39-2. 20 (2H, m), 2. .04-1. 79 (4H, m), 1. 72-1. 60 (1H, m), 1. 50-1. 18 (3H, m)	TH NUR (6) PPPP	

MC 872 (H+H)	Purity >90% (NMR)		Example No. 249	
		3.004££, D&SO-46 8. 02 (1H, 4], =1. 5Hz), 8. 11 (1H, 4, J=1. 8Hz), 7. 9G-7. 81 (3H, m), 7. 9G-1(1H, a), 7. 61-7. 49 (6H, a)), 7. 108 (2H, 4], 7-8. 6 1+2), 5. 19 (2H, s), 4. 25 (1H, m), 2. 38-2. 17 (2H, m), 1. 96-1 .78 (4H, m), 1. 70-1. 55 (1H, m), 1. 46-1. 16 (3H, m), 1. 11 (9 H, s)	1H NAR(8) ppm	

EP 1 162 196 A1

Table 186

		_					
MS 558 (H+1)	Purity >90% (NMR)	tapota	Example No.	MS 493 (M+1)	Purity >90% (NMR)		Example No.
		ğ	254				253
•	7(3H, a)	300Mfs, DMSO-d6 8. 25(1H, a), 8. 02(1H, d,]=8, 4, 1 71s, 7, 90(1H, dd,]=8, 4, 1 71s, 7, 80-7, 71(2H, m), 7, 30(2H, d,]=8, 71s), 7, 33(2H, d,]=8, 71s), 7, 25(2H, d,]=8, 71s), 5, 46(2H, s), 4, 78(2H, s), 4, 31(1H, m), 2, 39-2, 1, 9(4H, m), 2, 30-1, 1, 5(4H, m), 1, 50-1, 1	IN NMR(6).), 1. 43-1, 18(3H, m)	300kHs, DMSO-d6 8. 21(1H, d., J=1, 5Hz), 7. 93(1H, d., J=8, 7Hz), 7. 85(1H, dd .]=8, 41, 5Hz), 7. 54-7, 47(2H, ch), 7. 40-7, 24(6H, ch), 7. 16(1H, d., J=3, 6Hz), 7. 11-7. 105(1H, d., J=3, 6Hz), 7. 11-7. 105(1H, d., J=3, 6Hz), 7. 11-7. 105(1H, d., J=3, 6Hz), 7. 19-3, 6Hz), 5. 26(2H, g.), 4. 96(1H. g.) 12. 32-2, 13(2H, g.), 1, 58-1, 55(1H, g.) 172(4H, ch), 1, 68-1, 55(1H, g.)	1H NAR(8) ppm

MS 609 (H+1)	Purity >90% (NMR)		\ <u>\\</u> \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		ю.	Example No.
)	MR)	,	ð		>	252
		. 20 (2H, m), Z. 0), 1. 75-1. 64 (1 . 28 (3H, m)	7. 09 (1H, 1, J= (2H, 8), 4. 08 (1	1H, d, J=8. 6Hz) , J=8. 6, 1. 5Hz)	300MHz, DMSO-<	1H NMR(8) ppm
)1-1, 80(4H, E H, 四), 1, 51-1	7. 3Hz), 5. 43 H, m), 2. 40-2	7. 84 (1H, dd	46 1. 5Hz), 7. 96(8

Example No. 256	TH NAR (6) PDB
10	300kHz, DMS0-d6 12. 67 (1H, brs), 8. 23 (1H, s)
	6Hz), 7. 79(1H, dd, J=8, 7, 5, 6Hz), 7. 62-7, 41(7H, m), 6. 8
7	0(1H, dd, J=11. 9, 2. 3Hz), 6. 69(1H, dd, J=8. 1, 2. 1Hz), 5.
\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	20(2H, S), 3. 93(1H, brt, J=1 5. 3Hz), 2. 30-2. 11(2H, brm) 1. 88-1. 74(4H, brm), 1. 64-1
Furity >90% (NMR)	.58(1H, brm), 1.41-1.14(3H
SW 689 (H+1)	

	568 (H+1)	S
	Purity >90% (NMR)	Pur
1. 93-1. 78 (2H, m), 1. 73-1. 5 7 (1H, m), 1. 55-1. 16 (3H, m)	C	
8. 8Hz), 5. 54 (ZH, s), 4. 38 (1 H, m), 2. 74 (3H, s), 2. 40-2. 1	からなっ	
8. 34 (1f. 8), 6. 32 (1f. 6, Je8 . 8Hz), 8. 09–8. 03 (3H, m), 7. 83 (2H, d, J=8. 3Hz), 7. 79 (2H		
OME DASO-d6		
1H NAR(6) ppm	Example No. 255	Exa

Purity >90% (NMR)	***************************************	Example No. 260	MS 591 (N+1)	Purity >90% (NMR)		Example No. 259	MS 567(A+1)	Purity >80% (NMR)		Example No. 258	
m) 1. 50~1. 10 (3H, m)	300MHz, DMSO-d8 8. 93 (2H, d, J=6, 3Hz), 8. 35 (1H, s), 8. 26 (1H, d, J=8, 7Hz) 7. 80 (2H, m), 7. 86 (2H, 7. 35 (2H, d, J=8, 4Hz), 7. 24 (2 1. 35 (2H, d, J=8, 4Hz), 7. 26 (2H, s), 7. 35 (2H, d, J=6, 2Hz), 1. 90- 1. 75 (2H, m), 1. 70-1. 55 (1H, d)	IH NAR(6) ppm		2H, m), 1. 70-1. 55 (1H, m), 1. 50-1. 16 (3H, m)	300MHz, DMSO-66 8. 93 (2H, d, J=6, 6Hz), 8. 36 (1 11h. s); 8. 28 (1H, d, J=6, 7Hz) 8. 10-8. 03 (3H, m), 7. 85 (2H d, J=8, 7Hz), 7. 33 (2H, d, J= 8. 7Hz), 7. 23 (1H, s), 7. 23 (1 H, s), 6. 81 (1H, s), 5. 56 (2H, h, s), 4. 39 (1H, m), 2. 97, 2. 92 6H, s), 2. 40-2. 18 (2H, m), 2. 16-1. 95 (2H, m), 1. 50-1. 75 (IH NMR(6) ppm			300HHz, DMSO-d6 7.79 (1H, d., J=6, Th1), 7.56 (1H, d., J=7, EHs), 7.49 (2H, d., J=8, 6Hs), 7.42 (4H, s), 7.32 7.22 (3H, m), 7.09-7.03 (3H, m), 5.02 (2H, s), 4.46 (1H, m) 1, 3, 82 (3H, s), 1.95-1.83 (2 H, m), 1.75-1.44 (6H, m), 1.3 0-1.10 (2H, m), 0.89-0.71 (1 H, m)	IH NAR(6) ppm	

262

ö

8

Purity š

>90% (NMR) 564 (H+1)

ŧ

Example No.

266

1H NAR(8) ppm

K

Purity

>90% (NMR)

608 (H+1)

S

8

300Mrz, DMSO-d8 8. 30(14, d., J=1, SHz), 8. 25(1 18, d., J=3, 1Hz), 8. 03(1H, dd J=8, 7, 1. 5Hz), 7. 76-7, 96(1 3H, m), 7. 55-7. 49(5H, m), 7. 42(1H, d.), 27. 6Hz), 7. 23(2H 42(1H, d., J=7. 6Hz), 7. 23(2H, m), 9. 14 35(1H, m), 3. 0. 13Hz, 9. 2. 9 7(3H, 8), 2. 37-2. 20(2H, m), 2. 9 2. 09-1. 97(2H, m), 1. 94-1. 9 1(2H, m), 1. 72-1. 30(1H, m), 1. 50-1. 21(3H, m)

13

Example

NO.

265

1H NMR(6) ppm

8

Purity 3

>90% (NMR) 586, 588 (H+1)

3

300MHz, DMSO-d6
8. 23(H, d, J=1, OHz), 7, 92(
1H, dd, J=8, 7, 1, OHz), 7, 87(
1H, dd, J=8, 7, 1, OHz), 7, 60(2H, d, J=8, 7Hz), 7, 60(2H, d, J=8, 7Hz), 7, 20(1H, d, J=8, 7Hz), 7, 20(1H, d, J=8, 3Hz), 7, 21(1H, d, J=8, 2Hz), 7, 11(2H, d, J=8, 7Hz), 7, 106(1H, d, J=8, 7Hz), 7, 20(1H, d, J=8, 7Hz), 7, 36(1H, d, J=8, 36(2H, m), 2, 60-2, 40(2H, m), 2, 60-2, 20(2H, m), 2, 20-2, 20(2H, m)

õ

Example No.

264

1H NMR(&) ppm

MS 580 (H+1)	Furity > 90% (NMR)		Example No. 262	
) 58 (1H, m), 1. 62-1. 20 (3H, m	300Mz, DMSO-d8 8. 29 (1H, d, J=1, Shz), 8. 26 (1H, d, J=9, 0Hz), 8. 19 (1H, d, J=9, 0Hz), 8. 13 (1H, brs), 8. 08-7, 96 (2H, b), 7. 73 (2H, d, J=9, 0Hz), 7. 57-74, 316H, b), 7. 72 (2H, d, J=9, 0Hz), 5. 14 (2H, s), 4. 38 (1H, b), 2. 38-2 (2H, s), 4. 38 (1H, b), 2. 38-2 (2H, s), 4. 36 (1H, b), 1. 73-1	1H NAR(6) ppm	

병

ä

벊

MS 567 (I+1)	Purity >90% (NMR)		Example No.
	(4H, m), 1. 72-1. 57 (1H, m), 1 . 50-1. 18 (3H, m)	8. 22 (1H, d, J=7. 8Hz), 7. 85 (1H, d, J=6. 7Hz), 7. 65 (2H, d, J=6. 7Hz), 7. 65 (2H, d, J=9. 0H), 7. 51-7. 38 (6H, m), 7. 29 (1H, d, J=6. 3Hz), 7. 23 (1H, d, J=3. 0Hz), 7. 06 (2H, d, J=8. 6.3. 0Hz), 8. 05 (2H, e), 4. 41 -4. 26 (1H, e), 3. 33 (3H, e), 2. 40-2. 20 (2H, m), 2. 03-1. 78	261. 1H NMR(8) ppm

EP 1 162 196 A1

EP 1 182 198 A1

2

Purity

>90% (NMR)

300MHz, DMSO-d6 8. 27(1H, d, J=1, SHz), 8. 20(1H, dd J=8, 6, 1, SHz), 7. 82(2H, d, J=8, 61, SHz), 7. 82(2H, d, J=8, 2Hz), 7. 76-7. 65(6H, m) 7. 56(1H, dd, J=7, 91, 18Hz), 7. 47(1H, d, J=7, SHz), 7. 20 (2H, d, J=8, GHz), 5. 16(2H, s)), 4. 32(1H, m), 3. 02(3H, s), 2. 98 (3H, s), 2. 38-2. 19 (2H, s), m), 2. 07-1. 55 (2H, m), 1. 93-1, 80 (2H, m), 1. 72-1, 68 (1H, m), m), 1. 52-1. 18 (3H, m)

642 (N+1)

S

Example No.

269

Purity

>90% (NMR)

626 (M+1)

5

S

Purity

>90% (NMR)

t

300MHz, DMSO-d6, 8, 30(1)H, 8, 67-8, 59(1)H, m), 8, 02-8, 20(2)H, m), 8, 02-7, 29(2)H, m), 7, 65(1)H, t, 1-8, 18(1)H, dd, 1-12.0, 2, 2)Hz), 7, 18(1)H, dd, 1-12.0, 2, 2)Hz), 7, 16(1)H, dd, 1-12.0, 2, 2)Hz), 5, 14(2)H, m), 4, 09(1)H, m), 2, 8, 2(3)H, d, 1-4, 5)Hz), 2, 34-2, 1, 2(4)H, m), 1, 99-1, 79(4)H, m), 1, 1-1, 59(1)H, m), 1, 49-1, 2, 1, 1, 1-1, 59(1)H, m), 1, 49-1, 2, 1, 1, 1, m)

612 ()+1)

Example No.

268

IH MAR(8) ppm

Purity

>90% (NMR)

620 (H+1)

20

SW	Purity	<i>\</i>	Example No.	
652 (M+1)	>9.0% (NMR)		No.	
		^	271	
m)	1, 59(1H, m), 1, 50-1, 19(3H,	300kHs, DMSO-d6 8.29 (i.H. d, J=1. EHs), 8. 24 (IH. d, J=8. 7Hs), 8. 07-7. 98 (IH. d, J=8. 7Hs), 8. 07-7. 98 (3H, m), 7. 80-7. 88 (GH, m), 7. 56 (IH, dd, J=8. 0, 1. Shz), 7. 47 (IH, d, J=8. 0Hs), 7. 21 (2H d, J=8. 4Hz), 5. 18 (2H, s), 4. 3. (IH, m), 3. 27 (3H, s), 3. 80 2. (3H, s), 2. 80 (3H, s), 2. 38- 2. 18 (2H, m), 2. 10-1. 95 (2H,	1H NMR(8) ppm	

N.	שי		Į.	l
S	Purity		Example No.	
598 (H+1)	>90% (NMR)		٥.	
+1)	NMR)	.60	270	
		300kHz, DaSO-46 8.24(IH, d., J=1, 4kz), 8. 119(1, h, cr) 1. H, d., J=1, 8kz), 8. 11(IH, br) 8. 02-7, 85(3H, m), 7. 60 7. 44(7H, m), 7. 10(IH, dd, J= 12. 0. Z. 1kz), 6. 98(IH, dd, J= 12. 0. Z. 1kz), 5. 11(2H, s), 3 8. 4. 2. 1kz), 5. 11(2H, s), 3 1. 98(1H, m), 2. 30-2. 12(2H, m) 1. 91-1. 73(4H, m), 1. 71-1 58(1H, m), 1. 45-1. 15(3H, m)	1H NUR(6) ppm	

õ

EP 1 162 198 A1

Example No.

267

1H NMR(8)

300MHz, DMSO-d6 8.34(2H, m), 8.03(1H, d, J=8 8.34(2H, m), 7.77, 86(3H, m), 7. 54-7, 40 (4H, m), 7.33(2H, d, J=9.0 Hz), 5.18(2H, s), 4.36(1H, m), 3.01(3H, s), 2.97(3H, s), 3.01(3H, s), 2.97(3H, s), 2.40-2.20(2H, m), 2.11-1.9 7(2H, m), 1.93-1.81(2H, m), 7.1-1.60(1H, m), 1.50-1.2 1(3H, m) Table 191

MS 601 (H+1)	Purity about 8 0% (NMR)	.060.	Example No.
		300482, aixture 8, 35, 8, 3 0(2H, b), 7, 49 (2H, 2H, d, J=8, 2H, d, J=8, 1=8, 4Hz), 7, 207(1H, (2H, s), 4, H, s), 3, 7	. ~ I
		ure of cis and trans ,8.34(1H, s), 8.15–8.1 ,8.7.79–7.70(3H, s), (2M, d, J=8, THz), 7.44(,J=8, THz), 7.34(1H, d), ,J=8, THz), 7.31(1H, d), Hz), 7.25–7.19(2H, m) 7(1H, d, J=8, 5Hz), 5.08 ,3.70–1.90(8H, m)) ppm

MS · 645 Qt+ 1)	Purity >90% (NMR)		Example No. 273
	H, brm), 1. 49-1. 22 (3H, brm)	300kHs, DMSO-d6 8. 30(iH, s), 8. 27(iH, d, J=8 8. 7k±), 8. 05(iH, d, J=8. 7k±) 7. 77-7. 67(3H, m), 7. 58-7. 48(6H, m), 7. 22(2H, d, J=8. 4 46), 5. 18(2H, s), 4. 355(iH, b rt, J=9. 8k±), 3. 06-2. 88(12 H, brm), 2. 38-2. 20(2H, brm), 1. 90- 1. 80(2H, brm), 1. 70-1. 60(1	IH NAR(8) ppm

Table 193

EP 1 162 196 A1 Table 194

		50	Ġ	å	t	8	2	8	ŭ	6	Ch.
MS 635(M+1)	Purity >90% (NMR)	.⇔	to g	Example No. 278	Purity >90% (NMR) MS 61904+1)		Example No. 277	WS 603(H+1)	0		Example No. 276.
	brm), 2. 91-2. 70(2H, . 28-2. 11(2H, brm)	7. 22 (1H, d, j-2, 6H-2), 7. 13- 6. 92 (3H, m) 5. 05 (2H, s), 4. 6. 7 (1H, brt, j=14. 2H-2), 3. 57 -3. 40 (2H, brm), 3. 20-3, 05 (E, DMSO-d6 1H, bre), 8. 27 (1H 1nd7. 74 (2H, ABq, J 1. 58 (1H, t, J=8. 6H d7. 43 (4H, A' B' 6 7. 3 (1H, A' B' 6	IN MAR(6) ppm	s), 4.47and4.34(total IH, each brs), 3.83(3H, s), 3.12-1.7 6(8H, m)	H. each e), 7,94-7,87(IH, m), 7.60- 7.41(5H, m), 7.31(IH, d, J-8 .5Hz), 7.23-7,21(IH, m), 7. 12-7,05(2H, m), 7.00-6.88(IH, m), 5.06and5.05(total 2H, each	1H NMR(&) ppm 300Whz, DMSO-d6 cls and trans mixture 8.28end8.24(total	. u.w, 6. 21 ° . 10 (21, 01m)	1, 7, 09-6, 95(3H, m) H, s), 4, 11(1H, brt, L), 3, 84(3H, s), 2, 8 2H, brm), 2, 50-2, 32	300MHz, DMSO-d6 8, 25(1H, s), 7, 93end 7, 87(2 8, 25(1H, s), 7, 55(1H, t) 1, ABq, J-9, 1Hz), 7, 55(1H, t) 1, J=8, 6Hz), 7, 48end 7, 42(4H 4, J=8, 5Hz), 7, 24(1H, d, J=2,	IH NAR(6) ppm

å

Example No.

8

ĸ Purity g

2

8

S Purity ä

õ

Purity ₹ Example No. >90% (NMR) 644 (H+1) 279 300MHz, DMSO-d6 8. 30(1H, a), 8. 23(1H, d, J=8 7.71z), 8. 66-8. 00(2H, m), 7. 83(1H, dd, J=9. 0, 1. 8Hz), 7. 83(1H, dd, J=9. 4Hz), 7. 64(1H 7. 1(2H, d, J=9. 4Hz), 7. 64(4H 1. 1, 1. IH NIR(6) ppm

EP 1 162 196 A1

EP 1 162 196 A1

Example No. Example No. >90% (NMR) >90% (NMR) >90% (NMR) 630 (X+1) 580 (X+1) 284 283 282 300MHz, DUSO-d6 8, 30(1H, s), 8, 14(1H, d, J=8 8, 71m), 7, 97(1H, d, J=8, 71s) 7, 16(1H, d), 7, 16(1H, d), 7, 16(1H, d), 12, 2, 42, 2, 21s), 7, 03(1 H, dd, J=12, 4, 2, 21s), 5, 15(2 H, s), 4, 15(1H, s), 3, 54-3, 1 6(4H, s), 2, 33-2, 13(2H, s), 1, 70-1, 0 1, 97-1, 79(4H, s), 1, 70-1, 0 300MHz, DMS0-d6 30,03 (H, s), 8, 33 (IH, s), 8, 25 (IH, d), 96, 712), 8, 66 (I H, d, J=9, 0Hz), 7, 74 (2H, d, J) -9, 0Hz), 7, 51-7, 42 (6H, m), 7, 37-7, 30 (2H, m), 7, 22 (2H, d), 7, 22 (2H, d), 7, 30 (3H, s), 2, 40 37 (IH, m), 3, 06 (3H, s), 2, 40 -2, 18 (2H, m), 2, 15-1, 95 (2H, m), 1, 90-1, 80 (2H, m), 1, 75 -1, 65 (IH, m), 1, 1, 50-1, 20 (3H, m), m) 300Miz, DMSO-d6 8. 36(1H, s), 8. 35 (1H, d, J=9 3Mz), 8. 09(1H, d, J=9, 3Hz) 7. 78(2H, d, J=8, 7Hz), 7. 48 -7. 26(9H, m), 5. 09(2H, s), 4 -7. 26(9H, m), 5. 09(2H, s), 2. 4 0-2. 15(2H, m), 2. 10-1, 95(2 H,m), 1. 90-1, 75(2H, m), 1. 7 0-1, 55(1H, m), 1. 50-1, 20(3 H,m) IH NMR(8) ppm IH NAR(6) 1H NMR(8) ppm

Purity 5

654 (H+1)

400MHz, DNSO-d8 8. 29(1H, a), 8. 18(1H, d, J=8 6Hz), 7. 88(1H, d, J=8, 8Hz), 7. 72(1H, s), 7. 64(1H, T, J= 8. 8Hz), 7. 57-7, 43(6H, m), 7 1. 8(1H, dd, J=12, 12, 2 1Hz), 7. 03(1H, d, J=12, 12, 2 1Hz), 5. 12 (2H, a), 4. 15-4, 01(1H, m), 3 .75-3, 33(8H, m), 2. 31-2, 14 (2H, m), 1. 58(1H, m), 1. 47-1, 21

1H NUR(8) ppm

Purity

>90% (NMR)

652()(+1)

Example No.

	27.03(11, d. J=8.4Hz), 5.15(28, s), 4.07(11, m), 3.58-3, 41(41, m), 2.34-2, 13(21, m), 1.97-1, 77(81, m), 1.71-1, 58(111, m), 1.49-1.18(31, m)	300Hz, DMSO-d6 8. 29(1H, s), 8. 13(1H, d, J=8 . 0Hz), 7. 97(1H, d, J=8. 4Hz) . 7. 83(1H, s), 7. 68-7. 41(7H . m), 7. 17(1H, d, J=12. 0Hz),	287 IH NMR(6) ppm		1-3, 23(2H, a), 4, 07(1H, a), 3, 7 1-3, 23(2H, a), 1, 98-1, 71(4 H, a), 1, 71-1, 18(10H, a)	300Hz, MSO-46 8. 29 (1)4, 9), 8, 13 (14, d, J=8, 71x), 7, 97 (14, dd, J=8, 7, 1, 44x), 7, 69-7, 40 (81, m), 7, 16 (11, dd, J=12, 0, 2, 24x), 5, 02 (14, dd, J=8, 4, 2, 24x), 5	286 IH NAR (8) ppm	0, 1. 18 (6H, each s)	7.06(1H, dd, J=8.4, 2 5.13(2H, s), 4.22-4. m), 2.34-2.13(2H, m) -1.78(4H, m), 1.72-1.	8. 37 (H, d, J-7, 3Hz), 8. 30 (H, s), 8. 19-8. 12 (2H, m), 8. 02-7. 95 (2H, m), 7. 65 (1H, t) J-8. 4Hz), 7. 56-7. 43 (5H, m) J-1. 81 (1H, dd, J-9, 2, 0, 1, 8Hz)	285 IH NUR (6) ppm
8	3 0	ŧ	\$		t	સ	25	8	ä		
MS 668 0(+1)	Service Course		Example No. 290	MS 6820+1)	0		Example No. 289	#S 642(#+1)	٠,١		Example No. 288

IH MAR(6) ppm

400M/a, DMSO-d6
8.28(1H, s), 8.11(1H, d, J=8
9.82), 7.96(1H, d, J=8, 9Hx)
7.68(1H, s), 7.62(1H, t, J=
8.2Hx), 7.55-7.41(6H, m), 7
15(1H, d, J=8.4Hx), 6.14(2H, s)
4.12-3,13(6H, m), 2.30-1.

Purity

>90% (NMR)

666 (K+1)

5

Example No.

Purity

>90% (NMR)

640 (H+1)

S

Example No.

EP 1 162 196 A1

IH NMR(6) ppm

300HH₃, DMSO-d6 8. 62(1N, m), 8. 31(1H, s), 8. 22-8, 14(2H, m), 6. 99(2H, d), 6. 99(2H, d), 7. 99(2H, d), 7. 197, 7 H₂), 7. 66(1H, m), 7. 19 (1H, dd, 1–8, 7, 2. 2H₂), 5. 14 (2H, s), 4. 11(1H, m), 3. 67–3 (2H, s), 3. 45–3, 30(2H, m), 2. 00–1), 2. 17–2. 12(2H, m), 2. 00–1 .76(4H, m), 1. 70–1. 58(1H, m), 1. 17(3H, m)

Table 198

Purity

>90% (NMR)

	-	

š Purity Example No. S Purity Example No. >90% (NMR) >90% (NMR) 581 (1+1) 581 (H+1) 296 295 300MHz. DNSO-d6 8. 21(H, d, J=1, SHz), 7, 98(1H, d, J=1, SHz), 7, 97-7, 91(1H, d, J=1, SHz), 7, 97-7, 91(2H, ω), 7, 84 (H, d, J=2, 1Hz), 1, 770(H, d, J=7, SHz), 7, 60 -7, 54 (4H, ω), 7, 43 (H, d, J= 8, 4Hz), 7, 99 (2H, d, J=8, THz), 2, 50 (2H, s), 4, 25 (H, brt 1, J=14, SHz), 2, 36-2, 18 (2H, brt), 1, 36-1, 79 (4H, brt), 1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36-1, 36 1H MMR (6) ppm 1H NAR(8) ppm 8. 25and8. 04(2 Ha). 7. 74(IH. 5 J=8. 7Ha). 7. 48-7.3 I(2H. d. J=8. 7H I). 4. 46(2H s). J=14. 8H2). 3 37-2. 17(2H, b 52-1. 20(3H, bm), 1.

8

ä

ä

2

Purity S Example No. >90% (NMR) 567 (H+1) 1H NMR(8) ppm . 25and8. 03 (2 22) 7. 73 (1H, 5 J=8. 6Hz) 7. 5 0, 2. 3Hz) 7. 4 (1H, d. J=8. 0H J=8. 6Hz) 8. 5 5 (2H, s) 4. 35 5 (2H, s) 7. 2 2. 09-1. 2 37-2 2. 09-1. 2 5 (2H 5 (2H, bra) 1. 50-1

EP 1 162 196 A1

Purity 돐

1H NAR(6) ppm

Example No.

EP 1 162 196 A1

MS 562(H+1)	Purity >90% (NMR)		Example No. 299
		300KHz, DMSQ-46 8. 43-8, (A1, m), 8. 07-7. 9 4 (2H, m), 7. 72(H, d, J=8. 6H 2), 7. 62-7, 49 (5H, m), 7. 23 (2H, d, J=8. 6Hz), 5. 16 (2H, s) 4. 34 (1H, m), 2. 39-2. 22 (2H m), 2. 10-1. 96 (2H, m), 1. 93 -1. 80 (2H, m), 1. 71-1. 58 (1H m), 1. 49-1. 19 (3H, m)	IH NAKR (8) ppm

MS 599 (M+1)	Purity >90% (NMR)		Example No. 298	
	47-1. 17 (3H, brm).	300MHs, DMSO-46 8. 22(1H, s), 8. 01(1H, s), 7. 95and7, 86(2H, Afa, J=8, 6Hz,), 7. 79(1H, d, J=7, 8Hz), 7. 5 8(3H, t, J=7, 5Hz), 7. 53(4H, s), 7. 13(2H, d, 8, 7Hz), 5. (15, 15, 15, 15, 15, 15, 15, 15, 15, 15,	IH NMR(8) ppm	

2

Purity

>90% (NMR) ·

300HHz, DMSO-d6 8. 22(1H, s), 7. 95(1H, d, J=8 8. 7(1H), 7. 87(1H, d, J=1.5Hz 9. 0Hz), 7. 62(4H, d, J=3.4H 2), 7. 55(1H, 1, J=9.0Hz), 7. 20(1H 44(4H, d, J=8.1Hz), 7. 20(1H), 7. 11 (1H, dd, J=2.1Hz, 12.0Hz), 7. 11 (1H, dd, J=2.1Hz, 13. 7Hz), 6. 86(1H, s), 3. 94(1H, m), 2. 96 2. 88(1ZH, s), 2. 35-2. 00(2 H, m), 1. 95-1. 70(4H, m), 1. 6 5-1. 50(1H, m), 1. 45-1. 10(3 H, m)

663 (H+L)

Example No.

302

IH NAR (6) ppm

90% (NMR)	5	J)'
. 47-1. 20 (3H, brm)	5(2H, s), 4.26(1H, brt, J=13 .0Hz), 2.54(3H, s), 2.38-2. 20(2H, brm), 1.97-1.80(4H, brm), 1.71-1.59(1H, brm), 1	Hs), 7.60-7.55(3H, m), 7.49 and7.45(4H, A'B'q, J=8.3Hz), 7.12(2H, d, J=8, 7Hz), 5.0	300MHz, DMSO-d6 .12.7(1H, brs), 8.21(1H, s), 7.94and7.85(2H, ABq, J=8.6

Example No.

301

1H NAR(8) ppm

8

Purity

>90% (NMR)

523 (Y+1)

300

IH NAR(6) ppm

Example No.

SH Purity

>90% (NMR)

532 (M+1)

268

ô

Example No.

308

IH NAR(6) ppm

ä

Purity

>90% (NMR)

617(出+1)

PG.

Example No.

307

IH NUR(8) ppm

8

ĸ

577 (H+1)

Purity

>90% (NMR)

300MHz, DISO-dB 21 (H, 8) 12. 94 (JH, brs), 8, 21 (JH, 8) 7. 98-7. 84 (SH, m), 7. 56 (2H d, J=8, 7Hz), 7. 54 (ZH, d, J= 7. 84Hz), 7. 34 (JH, d, J=2, 4Hz), 7. 13-7. 06 (3H, m), 5. 06 (ZH, s) 4. 26 (JH, brt, J=12, 7Hz), 3. 84 (3H, s), 2. 35-2. 17 (ZH, b) rm), 1. 99-1. 80 (ZH, brm), 1. 77-11. 59 (JH, brm), 1. 47-1. 1

306 306 TH NAW (6) DDD

Example No.

EP 1 162 196 A1

EP 1 162 196 A1

268

3 Purity

>90% (NMR)

552 (M+1)

MS 616 (H+1)	Purity > 90% (NMR)) Q.		101	Example No. 310
	. 49-1. 18 (3H, brm), 1	n), 4. 66 (2H, 8), 4. 23 (1H, 6); t, J=11.8Hz), 3. 76 (3H, s), 2 t, J=2. 20 (2H, brm), 2. 04-1. 93 (2H, brm), 1. 89-1. 79 (2H,	71 (4H, A'B' q, J=8, 0Hz), 7, 4 3 (2H, d, J=7, 8Hz), 7, 15 (1K, d, J=8, 7Hz), 7, 07-7, 02 (4H,	300MHz. DMSO-d6 8.33(1H, s), 8.09and7.95(2 H ARG [58 7Hz) 7 87and7	IH NUR(8) ppm

_			
SM	Purity	ک	Example No.
	>90% (NMR)		No.
	(NMR)	.06.	309
	1. 72-1. 59 (1 . 17 (3H, brm)	300Mfz, DI 8, 33 (H, H, ABq, JH 69 (4H, A' 6 (2H, d, J' 6 (3H, m), 6 (3H, m), 2. 9Hz), 3, 2. 9Hz), 3, 2. 1 (2H, b)	IH NAR(6) ppm
	1) (1H, bra), 1. 49	NSO-d8 1, 8. 15and 7. 99 (2 8. 942), 7. 84and 7. 8. 942), 7. 22-7. 17. 01-6. 88 (24 m.), 7. 01-6. 88 (24 m.), 23 (24, A B Cq.) J-1 78 (34, 9. 3, 9-2, 9-2, 20, 20, 20, 20, 20, 20, 20, 20, 20, 2) ppm
<u></u>	<u>-</u> -	~ 262, 14.7 <u>6</u>	

Table 205

EP 1 162 198 A1

-				_	7.	-	_		-	-		
MS 628(H+1)	Purity >90% (NMR)	i, of o	Example 10.	Purity > 90% (NMR) MS 622041)		Example No. 313	609 (H+1)	rity >9			Example No. 312	
.81-1,71 (1H, m), 1.46-1.23 (3H, m)	1, d, J=4. 8Hz), 2. 32-2. 1, m), 2. 03-1, 87 (4H, m)	. 7. 66 (1H, d, J-7. 24 (6H, m), 6 -8. 6, 2. 6Hz), 6 -11. 6Hz), 2. 2H 0 (1H, m), 4. 99 (1H, m), 3. 95 (3H	1H NAR(6) DDB 300MHs, CDC13 8.48(1H, d, J=1.4Hz), 8.05(1H, d, J=1.8Hs), 8.98(1H, d, 1-8.6Hz), 7.82(1H, d, J=7.9	05 (2H, m), 2. 00-1. 50 (5H, m) , 1. 45-1. 10 (3H, m)), 7.89(1H, d, J=9.9Hz), 1-7.55(4H, m), 7.43(2H, 1-7.735(1H, t, J=0 1-7.732), 7.34(1H, t, J=0 1-7.724(1H, d, J=12.0Hz), 8.9 14(1H, d, J=8.6Hz), 8.9 H, 9), 3.96(1H, m), 2.35-	1H NSR(6) ppm 300MHz, DMSO-d6 8. 89(1H, brs), 8. 63(1H, brs) 9. 8. 24(1H, s), 8. 11(1H, d, J, J, J, S) 97. 8Hz), 7. 99(1H, d, I=8, 8H)		-1	12. 0Hs), 7, 13 (1H, d, J=8. 6H z), 6. 97 (1H, s), 3. 92 (1H, m) , 2. 35-2. 00 (2H, m), 1. 95-1. 70 (4H, m), 1. 65-1. 55 (1H, m) , 1. 50-1. 05 (3H, m)	. DMSO-d5 H, 8), 8. 12 (1H, d, J 8. 00-7. 84 (5H, m), d, J=8. 4Hz), 7. 56 (6Hz), 7. 23 (1H, d,	1H MHR(8) pi	

	412 (H+1)	SW
	Purity >90% (NMR)	A.
300MBz, DASO-66 8.23(IH, s), 7. 76(IH, d, J=8 8.23(II, s), 7. 76(IH, d, J=8, 8Hz) 7. 751-7. 32(7H, m), 7. 17(2H 7. 51-7. 32(7H, m), 7. 17(2H 8. 74z), 6. 55(IH, s), 5 18(2H, s), 4. 75(IH, m), 2. 3 5-2. 12(2H, m), 2. 10-1. 85(4 11, m), 1. 80-1. 56(2H, m)		3
1H NMR(8) ppm	Example No. 503	Œ

EP 1 162 198 A1

Table 208

272

Purity

>90% (NMR)

676 (H+1)

5 Purity Example No. Example No. >90% (NMR) 670 (M+1) 316 317 1H NHR (8) pps 1H NAR(8) ppm

t

300MHs, DASO-d6 9. 23 (III, t., 1-6, 1Mhs), 8. 29 (III, a), 8. 27-8. 22 (ZH, as), 8. 03 (ZH, d, 1-7, 9Hz), 7. 58-7. 48 (Eh, a) 7. 24 (4H, d, 1-4, 4Hz), 7. 28-7. 22 (3H, a), 5. 15 (ZH, a), 4. 52 (ZH, d) t. 1-8, 9Hs), 4. 35 (III, br t. 1-12, 1Hs), 2. 37-2. 18 (ZH, a) 2. 08-1. 95 (ZH, a), 1. 91-1. 79 (ZH, a), 1. 27-1. 59 (IH, a), 1. 47-1. 19 (SH, a)

ષ્ઠ

ä

Purity

>90% (NMR)

538 (X+1)

졼

315 300HHz, DMSO-46 8, 84(ZH, d., Jr.6, SH₂), 8, 28(1H 8), 8, 17mm² 99(ZH, AS₆, Jr.8 -7, 80(3H, a.), 7, 22 (1H, d., Jr.6, III NER(8) ppn

Example No.

8

Ġ

å

ä

3

8

至

EP 1 162 196 A1

EP 1 162 196 A1

Example

ĕ

321

IH NUR(8) ppm

Purity Example No. Purity Purity Example No. >90% (NMR) >90% (NMR) >90% (NMR) 671 (H+1) 671 (H+1) 663 (H+1) 323 322 300MH, MSD-dB
11. 19(1)k br
a), 8. 23and3, 02 (2
k, Ma, Jes, 0)tz), 7, 77 (1)k s), 7
7, 22and7, 22 (4)k le p, 1-8; 7k
17, 59and7, 48 (2)k, 8 g, 1-7;
9th), 7, 59and7, 48 (2)k, 8 g, 1-7;
9th), 7, 59and7, 14 (4)k, 8 g, 1-7;
9th), 7, 59and7, 14 (4)k, 8 g, 1-7;
9th), 7, 59and7, 14 (4)k, 8 g, 1-8;
18 (1)k, 1-8, 1-8;
18 (2)k, 1-8, 1-8;
19 (2)k, 1-8, 1-8;
19 (2)k, 1-8;
11 (2)k, 1-8;
11 (2)k, 1-8;
11 (2)k, 1-8;
12 (2)k, 1-8;
13 (2)k, 1-8;
14 (2)k, 1-8;
14 (2)k, 1-8;
15 (2)k, 1-8;
16 (2)k, 1-8;
17 (2)k, 1-8;
18 (2)k, 1-8;
18 (2)k, 1-8;
18 (2)k, 1-8;
18 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 (2)k, 1-8;
19 IH NAR(6) NER(6) Bdd gg

274

돐

MS 685 (H+1)	Purity >90% (NMR).	4	Example No. 32	
and the state of t	2H, m), 1, 91-1, 77 (2H, m), 1, 70-	300MH, 1850-d6 8.86(IH, I-7-6, 0Hz) 8.94and8 1.00 (4H, I-8-G, 1-8-G) Hz, R. 33 (IH 1.01 (4H, I-8-G, 1-8-G) Hz, R. 37 (IH 1.01 (1H, I-8-G, 1-8-G) Hz, T. 7-4an 1.01 (2H, A'B' G, 1-8-G) Hz) T. 7-4an 1.01 (2H, A'B' G, 1-8-G) Hz) T. 7-5and7 (8) (4H, A'B' G, 1-8-G) Hz) 1.53and7 (8) (4H, A'B' G, 1-8-G) Hz) T. 5-63and7 (8) (4H, A'B' G, 1-8-G) Hz) 1.50 (3Hz) 3.31 (2H, E, 1-8-G) Hz) 1.50 (3Hz) 3.31 (2H, E, 1-6-G) Hz)	325 IH NAR(8) ppm	

	662 (H+1)	SH
97.04\07.8)	ty >90% (NMR)	Purity
1, 3 12. 142) 3, 80 (14, br s), 2, 39-2, 18 (24, m), 2, 10-1, 9 8 (24, m), 1, 93-1, 57 (84, m), 1, 4	C	
.14 (2H, 8), 4.36 (1H, br	ص مړ	
28, A' B' q, J=8, 3Hz), 7, 74and7.	, 6 6 7 6	¥
28	J°	
Suche process		
TH NAME (&) BOTH	394	

Table 211

EP 1 162 196 A1

	683 (W+1)	SW	
	>90% (NMR)	Purity	3
. 51-2.05(2H, m), 1. 90-1. 70 (4H , m), 1. 65-1. 65 (1H, m), 1. 40-1. 10 (3H, m)	7		٠
8. 7Hz), 7. 70-7. 50 (5H, m), 7. 27 -7. 20 (3H, m), 7. 08 (1H, d, J=7. 8 Hz), 6. 90 (1H, s), 3. 93 (1H, s), 2		<u></u>	
13. 20-12. 60 (2H, brs), 8. 23 (1H .s), 7. 98 (2H, d, J=8. 6Hz), 7. 95 (1H, d, J=8. 7Hz), 7. 87 (1H, d, J=	, ż	₹	•
IH NAR(6) ppm	o. 327	Example No.	
	Table 212		•

EP 1 162 186 A1

Table 213

t			50			å		\$			u			3 6		8			8			ij.		;	5		•
2050	2049	2048	2047	2046	2045	2044	2049	2042	2041	2040	2039	2038	2037	2036	2095	2094	2093	2032	2031	2030	2029	2028	2027	2028	2025	2024	2023
5- (-Ac)	5- (-CO ₂ Me)	5- (-C0 ₂ Me)	4-(-CO ₂ H)	5- (-C0 ₃ H)	5- (-CF ₃)	5- (-CF ₃)	5- (-CE ₂)	5 (-Me)	5- (-Me)	5- (-Me)	4-(-NO ₁)	5~ (-NO ₂)	4- (-CN)	5-(-CN)	4- (-C1)	5- (-C1)	4-(-C1)	5-(-C1)	5-(-C1)	4-(-C1)	5-(-C1)	5-(-C1)	4-(-C1)	5-(-C1)	4-(-C1)	4-(-C1)	4-(-C1)
4- (-F)	4-(-C1)	4-(-8)	4-(-01)	4-(-8)	(\ \(\)	4- (-CO ₃ Me)	4- (-содн)	(گٹر))_	4-(-CO2Me)	4- (-CO ₂ H)	4- (-C1)	4-(-5)	4-(-C1)	4- (-F)	4_ (-8-m)	4- (-8-4)	4- (-SMo)	3-(-0Me)	4-(-CON (Me) 2)	4-(-CONH2)	(-LO)	4-(-CO ₂ Me)	4-(-co ₂ H)	4-(-CF ₃)	4-(-Me)	4-(-C1)	4- (-E)

Ex. No.

2001
2002
2003
2004
2006
2006
2008
2008
2009
2010
2011
2013

4-(-F)

6-(-F) 5-(-F)

5-(-F)

4-(-CE₃) 4-(-CO₂He)

4-(-C1) 4-(-Me)

4-(-F) 4-(-F) 3- (-F)

2-(-8)

3-(-P)

-H -H 5-(-F)

R' (-Me)

3- (-CF₃)

. -H

3-(-F)

2016 2017 2018 2019 2020

> 5- (-P) 5- (-F)

4-(-CON (Me);)
4-(-CMe)
4-(-SMe)

5- (-F) 5- (-F)

(OT)

4-(-CONH2)

5- (-F)

2015

2021

4-(-CL)

4- (-4-2)

86 A1

5 (10) 5 (10)

2066 2067

4-(-0%e)

4- (-NHAC)

4- (-NHMe)

2069

2065

5-(-1-O) 5-(-1-O)

4-(-C(-NR)NH2)

4-{-CON (Me);)

2060 2061 2062 2063 2084

5-(-1-(-)

; (10) ; (10)

4- (-CO₂Me)

4-(-co₂H)

4-(-CF₃) 4-(-Ac)

<u>5</u>(10)

 $4-(-\text{CONH}_2)$

2059

2051 2052 2058 2058 2054 2056 2057 2058

5_(-1-(-)

4-(-NO₂)

4-(-CN)

4- (-P)

5_(AO)

(<u>L</u>()

3			8			å			ò		;	:		1	ŝ		1	2		;	3		18		ō		•
2098	2097	2096	2095	2094	2093	2092	2091	2090	2089	2088	2087	2086	2085	2084	2083	2082	2081	2080	2079	2078	2077	2076	2075	2074	2073	2072	2071
5-(-CONH ₂)	5- (-conh2)	5- (-CONH ₂)	5- (-CONH ₂)	5- (-CONH ₂)	5-(-CONH ₂)	5- (-CONH ₂)	5-(-CONH2)	5-(-conh ₂)	6-(-CONH ₂)	6-(-CONH ₂)	6-(-CONH ₂)	4-(-CONH ₂).	4-(-CONH ₂)	4-(-conh _z)	3-(-CONH ₂)	3- (-CONH ₂)	3- (-CONH ₂)	5-(-con ₁₂)	5- (-CONH ₂)	5- (-CONH ₂)	5- (-CONH ₂)	5-(-CONH ₂)	₅_(♣ ○)	5_(┸Ѻ)	_{5_} (-1-(-))	(○₁ <u>+</u>)	_5_(○ -(○)
4-(-CO ₂ Me)	4-(-CO ₂ H)	4- (-Ac)	4-(-CF ₃)	2,6-d1-(-Me)	4-(-Me)	4-(-NO ₂)	4-(-CN)	3,5-d1-(-C1)	4-(-C1)	3-(-C1)	2-(-C1)	4-(-C1) .	3-(-C1)	2-(-C1)	4-(-C1)	3-(-C1)	2-(-C1)	3- (-C1)	2- (-C1)	2,3,4,5,6-penta-(-F)	4-(-F)	н-	$4-\left\{-\frac{9}{6}n(m),\right\}$	4_ (4- (4_()	4- (-SNe)

EP 1 162 196 A1

5- (-Ac)

-H

EP 1 162 196 A1

5- (-CONH₂)

4- (-1-)

4- (-SMe)

4-(-NHMe) 4-(-NHAC)

5- (-CONH2)

5- (-CONH₂)

2102

2101

2099 2100

5- (-CONH₂)

3,5-d1-(-CONH₂)

4- (-CONH₂)

2104 2105

5- (-CONH₂)
5- (-CONH₂)
5- (-CONH₂)
5- (-CONH₂)

3,4,5-tri-(-OMe)

4-(-CON(Me)₂)

4- (-OMe)

2106

2119

2116 2117 2118

4-(-CON (Me) 2) 5-(-CON (Me) 2)

5-(-CON(Me)₂)
5-(-CON(Me)₂)

2115

2114

2113

5- (-CONH2)

4 (-0

2110 2111 2112

5- (-CONH₂) 5- (-CONH₂)

2120

5-(-CON (Me) 2)

2121 2122 2123

> 4-(-CON (Me) 2) 5-(-CON (Me) 2)

5-(-CON (Me) 2)

4- (-CO2H)

4- (-CF3)

4-(-Ac)

4- (-NO2)

4-(-C1)

4- (-Me)

281

2171 2172 2173

5-(-NHAC) 5-(-NHAC)

5-(-NHAC)

4-{-CON (Me) 2}

4- (-F)

4- (-CONR2)

4-(-Ac)

4-(-C1)

4-(-F)

2168 2169 2170

5-(-NHMe)

2161 2162 2162 2183 2184 2164 2166 2166

5- (-OMe)

{-

4-(-C1)

4- (-F)

5-- (-NEMe)

5- (-OMe)

5- (-OMe)

5- (-OMe)

2159

5- (-OMe)

(- q l ()

4-(-CON(Me) 2)

4- (-OMe)

5- (-OMe)

2180

5- (-OMe)

4- (-1--)

4- (-SMe)

4-(-NHMe) 4-(-NHAC)

5- (-QMe)

2157 2158

2155 2156

5- (-OMe)

5- (-OMe)

2164

5- (-OMe)

(PO)

4- (-CO₂Me)

4-(-CO₂H)

4-(-CONH2)

2151 2152 2153

4,5-d1-(-OMe)

4- (-OMe)

EP 1 162 198 A1

EP 1 162 198 A1

2214 · 2215

S- (-10 mil)

2213

4-(-CON (Me) 2)

4- (-SMe)

4-(-CONH2)

4-(-Me) 3-(-CF₃)

4-

2212

2210 2211

S_ (-8-m)

2207

2,4-d1-(-C1)

4-(-C1)

4-(-F)

2209

2206

5- (-0-m) 5- (-0-m) 5- (-0-m) 5- (-0-m)

4-(-CON (Me) 2)

4-(-CONH₂)

4-(-CF₃)

4-(-C1)

2204

2202

2200

2198 2199

65	80		à	\$		놚		g			G	ষ্ট	1	c c		70		G.
2235	2234	2299	2232	2231	2290	2229	2228	2227	2226	2225	2224	2223	2222	2221	2220	2219	2218	2217
5- (\(\frac{1}{4} \frac{1}{4	5_(^A O**)	5- (Argan)	5- (- J O H)	5- (-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	5_ (~~~**)	_{5_} (~~\^\)	<u>, (~~))</u>	5-{-0-{CH ₂ },-OH}	5-(-0-(CH ₂) ₂ -OH)	5- {-g-10w}, }	5- {-{-(0m), }	5_ {-{-am}, }	5 {	5_ {-{-(un), }}	5- { f-+ (0m), }	5- {-8-4(10)+}	4- {	5_ {
4- (-C1)	4-(-Cl)	4- (-Cl)	4- (-01)	4- (-C1)	4- (-C1)	4~ (-C1)	4- (-C1)	4-(-C1)	4-(-C1)	4- (- f-u)	4- (-3-4)	4- (-SMe)	4-(-CON (Me) 2)	4- (-CONH ₂)	4-(-CF ₃)	4- (-Ma)	a- (-c1)	4- (-F)

EP 1 162 196 A1

4-(-CON (Me) 2)

4-(-F)

EP 1 162 196 A1

2249	2248	2247	2246	2245	2244	2243	2242	2241	2240	2289	2238	2237	2236
5 <u>-</u> ()	(الله)	<u>5_</u> (ŇѾ)	5_ (A Can)	([†] Q.)	<u>،</u> (أص)	ر ^ا ل (کرد))	5_(Å&)	<u>-</u> (^ا لکي)	<u>. ب</u> رځمي)	5- () Bay Jay ()	5- (*****)	ر ^{ال} ل _{تهرا})	("Oy)"
4-(-01)	4-(-C1)	4-(-C1)	4-(-C1)	4- (-C1)	4-(-Cl)	4-{-C1}	4~(-C1)	4-(-C1)	4-(-C1)	4-(-C1)	4- (-C1)	. 4-(-C1)	4-{-C1)

2254 2253 2252 2251 2250 4-(-C1) 4-(-01) 4-(-01) 4-(-C1) 4-(-01)

EP 1 162 198 A1

4-(-C1) 5-(-C1) 4-(-C1) 5- (-F)

4- (-Me) 4-(-C1)

4- (-E)

4-(-CF₃)

289

290

2271 2270 2269 2267 2268

4 (-2 4-(-1-1)

(PO)

2300 2301

5- (-co2Me) 5-- (-CO₃Me)

4-(-C1)

4-(-P)

4-(-CO2H) 5- (-созн)

4-(-C1)

4-(-F)

5- (-Ac)

5-(-Ac)

4-(-CL)

Ŧ

4-(-F)

4-(-F)

2297 2298

2296

· 5- (-CF₃)

2294 2295

2293

5- (-CF₃)

5- (-CE)

5- (-Me)

4-(-co₂H)

4- (-CO2Me)

(PQ)

2292

2265

2266

5- (-F) 5- (-F) 5-(-F) 5-(-F) S-(-P) 5- (-F) 5-(-F) 5-(-F) 5-(-F) 분 표 ᆈ Ø

5-(-8) 3-(-8)

4- (-QMe)

4- (-SMe)

5- (-P) 5-(-F) 2261 2262 2268 2268

5- (-Me) 5- (-NO2) 5- (-NO2)

5- (-Me)

4- (-CO2Me)

4-(-co2H) 4-(-01)

2286 2287 2288 2288 2289 2290 2291

5- (-CN)

2285

5-(-CL)

2284

5-(-C1)

4-(-8)

4-(-01)

4-(-F)

5-(-C1)

2282 2283

2281 2280 2279 2278 2277

5-(-C1).

4-(-0Me)

4- (-SMe)

6-(-C1) 5-(-C1)

4-(-CONH2) 4-(-CON (Ma) ;)

5-(-C1) 5-(-C1)

4- (-CO₃Me)

4- (-CO2H)

EP 1 162 196 A1

4-(-8-4)

4- (-SMe)

2319 2320

4-(-NHAC)

2313 2314 2315 2316 2316 2318 2312

; (10)

(LO)

4~ (-CO2Me)

4-(-Ac) 4-(-CO₂H) .

5-(±O)

5- (-PO)

4-(-C(=NH)NH2)

4-(-CON (Ne) 2)

4-(-CONH₂)

(المسلك)

4- (-0Xe)

4- (-NHM6)

2311

2910

<u>"</u>(上()

2309

§_(10)

2305 2306 2307 2307

(PO)

4-(-CF3)

4-(-C1) 4-(-NO₂) 4-(-Me)

5-(LO)

2304

1- (AO)

	80	,	ŧ	•		6		;	ŧ		8			2			18			š		70		G.
2347	2346	2345	2944	2943	2342	2341	2340	2339	2338	2337	2336	2336	2334	2999	2382	2991	2330	2329	2928	2327	2926	2926	2324	2923
5- (-CONH ₂)	5- (-conh ₂)	5~ (-CONH ₃)	5- (-conh ₂)	5-(-conh3)	S-(-CONH ₂)	5- (-conh ₂)	5- (-conh ₂)	5- (-CONH ₂)	5- (-conh ₂)	5- (-conh2)	(*RNOD-) ~5	5- (-conh³)	5- (-CONH2)	5- (-conh³)	5- (-CONH2)	5- (-CONH ₂)	5- (-CONH ₂)	5- (-conh ₁)	4- (-CONH ₂)	5- (-CONH ₂)	5- (-CONH ₂)	<u>, (LO)</u>	<u>5_(♣</u> ())	_(_ 1 _())
4-($\frac{4}{4}$ $\left(-\frac{9}{8}-\frac{1}{10}\right)$	4-(-SMe)	4-(-11-10)	4-(-NHAC)	4-(-NHMe)	(4- (-0Me)	4-(-C(=NH)NH ₂)	4-(-CON (Me) ₂)	4-(-CONH ₂)	<u>4</u> -(اللاص)	4-(-CO ₂ Me)	4- (-CO ₂ H)	4~ (-Ac)	4- (-CP ₃)	4-(-Me)	. 4-(-NO ₂)	4-(-CN)	4-(-C1)	4- (-F)	H	4-{	4_ (4- (-1-4-10)

EP 1 162 196 A1

2364 2365 2366 2367 2368 2368 2370 2371

5-(-CON (Me) 2)

5-(-CON(Me)2)

4-(-SMe)

5- (-CON (Me) 2)

4- (-1-1)

5-(-CON(Me)2)

4- (-NHAO)

4- (-NHMe)

5-(-CON (Me) 2)

5-{-CON(Me)2} 5-{-CON(Me)2}

4-(-------)

5-(-CON (Me)₂)

٤,

2362 2363

5-(-CON(Me)₂)

5-(-CON (Me) 2) 5-(-CON (Me) 2)

4-(-con(Me)₃)

4- (-CO2Me)

4-(-CO3H)

5- (-CON (Me) 2)

4-(-C(-NH) NH2)

4- (-QMe)

2359 2360 2361

2356 2357 2358

5- (-CON (Me) 2)

2349 2350 2351 2352 2353 2354 2355

5-(-CON(Me)₂) 5-(-CON(Me)₂)

4-(-NO₂)

4-(-CF₃)
4-(-Ac)

5-(-CON (Me) 2)

5-(-CON(Me) 2)

4- {-8-4(m), }

5- (-CON (Me) 1)

4-(-E) 4-(-C1) 4-(-CN)

EP 1 162 196 A1

2348

5- (-CONH₂)

4- (-8-m)

8		so		\$		8			ĸ		8			8			ষ্ট			ā.			õ		•
2398	2397	2996	2395	2394	2399	2392	2391	2390	2389	2988	2387	2386	2386	2384	2383	2382	2381	2380	2979	2378	2977	2976	2976	2974	2379
9-(~NHMe)	5-(-OMe)	5-(-OMe)	5- (-QKe)	5-(-OMe)	5-(-0Xe)	5-(-ome)	5- (-OMe)	(-OMO)	5-{-0Ne}	5~ (~OMe)	5- (-0Me)	5- (-OMe)	5- (-OMe)	5- (-OMe)	5- (-0%e)	5- (-OMe)	5- (-OMe)	5- (-0หุย)	5- (-0Me)	5- (-OMe)	5- (-0Me)	5- (-OMe)	(-OMe)	5- (~OMe)	5-(-con(Me) ₂)
4- (-F)	4-{-8-40, -}	4_ (-{\frac{1}{2}}-104)	4- (- -a)	4- ()	4-(-SMe)	4_ (-11-8-14)	4-(-NHAC)	4- (-NHMe)	4_(~~~ \\ ())	4-(-OMe)	4-(-C(-NH)NH ₂)	4-(-CON(Me) 2)	4-(-CONH ₂)	<u>(گٹر</u> ()	4-(-CO2Me)	4- (-CO ₂ H)	4-(-Ac)	4-(-CF ₃)	4- (-Me)	4-(-NO ₂)	4- (-CN)	4-(-C1)	4-(-F)	H	4- {- }+(sa), }

EP 1 162 196 A1

2415 2416 2417 2417 2418 2419 2420

2414

5- (-1 1-)

5- (-SNe) 5- (-SMe)

4-(-C1)

4- (-Me)

4-(-F)

4-(-CF3)

2412

2407 2408 2409 2410 2411

5- (-18 1)

(PO)

4-(-CO₂H)

S- (+ + + +)

2413

5-(100)

1

4- (-SMe)

4-

2422

5-(-SMe)
5-(-SMe)
5-(-SMe)

4-{-CON (Me) 2)

4- (-CONH2)

4-(-Ac)

	50		à		40	ŧ	ŧ	ક		8		*8		13		10		u,
2441	2440	2439	2438	2437	2436	2436	2434	2433	2432	2431	2430	2429	2428	2427	2426	2426	2424	2423
5_ (5- (-g-104)	5- (-g-nn)	5_ (-{ h m)	5- (-f- nh)	5- (- 8 -m)	5_ (-8-4.)	5_ (-8-4)	5- (-8-1)	5_ (- 1 4.)	5- ()	5_ (-{ -	5_ ()	5_ ()	5_ (-1-4-)	5_ ()	5- (-8-40)	5- (-1-4-)	5- ()
4-{-CON (Me) 2}	4- (-CONH ₂)	4-(-CP ₃)	4- (-Ne)	4-(-C1)	4 → (– E)	4-{-CON(Me) ₂ }	4-(-CONH ₂)	4- (-Ac)	4-(-CF ₃)	4- (-Me)	4-(-C1)	4-(-?)	4-{-CON(Me)2}	4-(-CONH ₂)	4-(-Ac)	4- (-CE ₃)	4-(-Me)	4- (-C1)

EP 1 162 198 A1

2400 2401 2402 2403 2404

> 5- (-NHAC) 5- (-NHAC)

2405 2406

5-(-1-1-1-)

5- (-NHAC) 5- (-NHAC)

4-(-CON (Me) 2)

4- (-CONH₂)

4-(-C1)

4-(-Ac)

4- (-CF3)

4-(-Me)

4-(-C1)

5- (-19-

2399

5- (-NHNe)

4-(-C1) 4-(-F)

EP 1 162 198 A1

4-(-F)

4-(-C1)

4-(----)

5- {-8-10m, }

5- (-1-00.)

2442

2445 2446 2447

5_ {-||-||-||(100), }

5- {-- (Ou), }

88		8			à		i	å		ŧ,			8			25			20			16			õ		•
2476	2476	2474	2473	2472	2471	2470	2469	2468	2467	2466	2465	2464	2463	2462	2461	2460	2459	2468	2457	2456	2455	2454	0 5	2			
3- (-NHMe)	₃₋ (-→,- ¹ -(○)	3- (-QMe)	3- (-C (-NH) NH2)	3-(-CON(Me)))	3-(-CON(Me);)	3-(-CON(Ma);)	3- (-CONH ₂)	3-(-CONH2)	3-(-CONH ₂)	₃_(^	3- (-co ₂ Me)	3- (-CO ₂ H)	3-(-Ac)	3- (-C£3)	3-(-xe)	3-(-NO ₂)	3- (-CN)	3,5-d1-(-c1)	3-(-C1)	2-(-F)	2~(-8)	2-(-F)	20		0		Tab
3~(-MHN-)	3-()	3- (-OMe)	3-{-C (-NH) NH ₂ }	3-(-C1)	3-(-8)	3-(-CON (Me) 2)	3-(-C1)	3- (-F)	3- (-con ₁)	ر (کار <u>)</u> (3- (-CO ₂ Me)	3-(-CO ₂ H)	3-(-Ac)	3- (-CE ₃)	3- (-Me)	3- (-NO ₂)	3- (-CN)	3,5-d1-(-C1)	3-(-C1)	4- (-£)	3-(-8)	2- (-F)	R'			\\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\	Table 215

2453

2451 2452 2448 2449 2450

4-(-CON(Me) 2)

4- (-CONH₂)

4-(-Me)

4- (-1-4)

4- (-SMe)

4 - (-5Ne)

EP 1 162 198 A1

298

	t	8			ä		ö		•	
2614	2519	2512	2511	2610	2509	2508	2507	2506	2505	2504
ر (مورسائي) مورسائي (مورسائي)	4_ (-1-44)	4- (4- (-4-11)	4- (-SMe)	4_ (-)	4-(-NHAC)	4-(-NHMB)	$_{4-}(\circ + 1)$	4- (-0Me)	4- (-OMe)
4- {-8-40a), }	4_ (-\frac{1}{12} ms.)	4_ ()	4- (- 1-1 -1-1)	4-(-SMe)	4_(- - - -)	4-(-NHAC)	4-(-NHMe)	<u>(</u> -(→- ^{al} l-())	3,4,5-tri-(-OMe)	4-(-OMe)

	ta:	29			ä		õ		,	•
2514	2519	2512	2511	2610	2509	2508	2507	2506	2605	2504
4- (-ق-اسی.)	4- (-f-m)	4- (- -lb)	4- (-l-m)	4-(-SMe)	4- (-#-g-m)	4-(-NHAC)	4- (-NHMe)	$q_{-}(\rightarrow a_1 h_1 \bigcirc)$	4~ (-OMe)	4-(-0Me)
4- {-8-0m, }	4_ (-8-m)	4_ ()	4- (- - - - - - - - - - - - - 	4-(-SMe)	4_ (4-(-NHAC)	4-(-NHMe)	(3,4,5-tri-(-OMe)	4-(-OMe)

4-(-CON (Me) 2) 4-(-CON(Me)2)

4-(-C(=NH) NH2)

3,5-d1-(-C1)

4-(-01)

4-(-CON (Me) 2)
4-(-CON (Me) 2)

2498 2497 2498 2499

4- (-CONH₂) 4- (-CONH₂) 4- (-CONH₂) 4- (-CONH₂)

2,3,4,5,6-penta-(-F)

4-(-B)

4-(-CL)

4- (-CON (Me) 2)

4-(-F)

2496 2494 2492 2493

(LO)

(LO)

4-(-CO2Me) 4- (-CO₂H)

4- (-CONH₂)

4- (-CO₃Me) 4-(-CO2H)

2489 2480 2491

4-(-CF₃)

4-(-Ac)

4- (-Me)

2,6-d1-(-Me)

4-(-CE)

4- (-Ac)

4- (-NO₂) 4- (-CN)

4-(-C1) 4- (-F)

4-(-Me)

2488 2486 2487 2485 2484 2483 2482 2481 2480 2479 2478 2477

4- (-NO₂)

3- (-Me)

4-(-CN) 3-(-C1) 3- {-1-4(10), }

3- (-8-17)

3- (-8-4) 3-(-9-1)

3-(-F)

3- (-18-2)

3- (-1-1)

3- (-1-1-1-)

3- (-#---)

3-(-SMe)

3- (-NHAC)

3- (-NHAC)

3- (-8Me)

300

EP 1 162 196 A1

2535 2536

3- (-NHMe)

3- (-NHAC)

3- (-NHMe)

2534

3_(~~~\frac{\frac{1}{2}}{2})

3-(------)

3-(-C(=NH)NH2)

3-(-C(=NH) NH2)

3-(-C1)

3-(-0Me)

3- (-OMe)

2526 2527 2528 2529 2530 2531 2532

3-(-CON (Me)2)

3-(-CONH₂)
3-(-CON(Me)₂)

3- (-CON (Me) 1)

3- (-C1)

3- (-F)

2518 2519 2520 2521 2522 2522 2523

> 3- (-Me) 3- (-CF))

3- (-Ac)

3- (-CN)

R -H 2-(-F) 3-(-C1)

3- (-NO₂)

3-(-NO2)

3- (-C1)

2516 2517

2625

(PO)

(PO)

3- (-CO3Me)

3-(-Me)
3-(-CF₃)
3-(-Ao)
3-(-CO₂H)

3- (-CONH₂)

3- (-F)

3- (-CO₂Me)

3- (-CO₂H)

3- (-CONH2)

Table 216

8		50	-		\$			ŧ			19		,	8			t.		8	20		õ		76		us.	
2563	2562	2661	2860	2559	2558	2557	2856	2655	2664	2553	2552	2661	2550	2549	2548	2547	2546	2545	2844	2543	2542	2641	2540	2539	2638	2537	
4- (-NHAC)	4-(-Nнме)	(\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	4- (-0%e)	4-(-C(=NH)NH2)	4- (-CON (Me) 2)	3- (-CON (Me) 2)	3- (-CON (Me) 2)	4-(-CONH ₂)	4-(-CONH ₂)	4-(-CONH2)	(O ⁺ C)	4-(-CO ₂ Me)	3- (-CO ₂ H)	4-(-Ac)	4-(-CF ₃)	(-Me)	4-(-NO ₂)	4-(-CN)	4-(-C1)	. 3-(-F)	3- {	3_ (-\$\$\text{\$\exiting{\$\text{\$\exititt{\$\text{\$\exititt{\$\text{\$\text{\$\texititt{\$\text{\$\texititt{\$\text{\$\text{\$\text{\$\text{\$\texititit{\$\text{\$\texititt{\$\text{\$\texititit{\$. 3_ (-100)	3_(-1/46)	3-(-SMe)	3_ (-#)	
4-(-NHAC)	4- (-NHMe)	(4-(-0Me)	4-(-C(=NH) NH ₂)	4-(-C1)	4- (-F)	4-{-CON(Me);	4-(-C1)	4-(-F)	4- (-CONH ₂)	(O+C)	4- (-CO ₂ Ma)	4-(-CO ₂ H)	4-(-Ac)	4-(-CF ₃)	4-(-Me)	4-(-NO ₂)	4-(-CN)	4-(-01)	. 4-(-8)	3- {-8-00, }	3_ (- -n)	3_(· 3- (-\$-\$)	3-(-SMe)	3- (-#)	

2

EP 1 162 196 A1

		8			ta ta			8			23			20			Ğ		70		
2587	2586	2685	2584	2589	2582	2581	2580	2579	2578	2577	2576	2676	2574	2573	2572	2671	2570	o.N			
3-Py	2-Py	3-Py	3-Py	ý-Py	3-Py	3-Py	3-PY	3-Py	3-Py	3-Py	3-Py	3−₽y	3-PY	3-Py	3-Ру	Åд~Ε	3-Py	ላል	(Table 217
4- (-CO2Me)	4- (-co ₂ H)	4-(-Ac)	4-(-CF ₃)	4-(-Ne)	4-(-C1)	4-(-F)	3-(-CON (Me) 2)	3-(-CONH ₂)	<u>3</u> -(♣○)	3-(-CO ₂ Me)	3- (-содн)	3- (-Ac)	3- (-CF ₃)	3- (-Me)	3-(-C1)	3~ (~F)	-8	₽'.	Py : pyridyl group	4.	217

EP 1 162 196 A1

4-Py 3-Py

4-(-CON(ME)₂)

3-Ру

2

2587 2588 2589 2590

2568 2567

4- (----)

2565 2566 2564

4- (-3Me)
4- (-3Me)
4- (-8-4)

4-(-SMe)

4-(-8-4)

Table 218

4-(-CON(Me)2)	11 3-Py	2611
4-(-CONH ₂)		2610
((L ())		2609
4-(-CO ₂ Me)	08 3-Py	2608
4-(-CO ₂ H)		2607
4-(-Ac)		2600
4-(-CF ₃)		2605
4- (-No)		2804
4-(-C1)		2603
4-(-F)		2802
3-(-CON (Me) 2)	01 3-PY	2601
3-(-CONH ₂)	00 3-Py	2600
(P)	99 3-2-у	2599
3- (-C0 ₂ Me)	98 3-EY	2598
3-(-CO ₂ H)	97 3-Py	2597
3- (-Ac)	96 3-Py	2596
3- (-CF ₃)	98 3-FY	2595
3-(-Me)		2594
3-(-C1)		2593
3- (- <u>F</u>)		2592
12		2591
72.		Ex.N
Py : pyridyl group	0	

ŭ

ä

[0301] Formulation Example is given in the following. This example is merely for the purpose of examplification and does not limit the invention.

EP 1 162 196 A1

Formulation Example

_	_		_	
ì	3	0	3	3
the same of the same	sodium carboxymethylcethulose	com starch	lactose	compound of Example 1
	44 9	150	809	100

(0303) The entire amounts of (a), (b) and (c) and 30 g of (d) are kneeded with water, dried in vacue and granulated. The obtained granules are mixed with 14 g of (d) and 1 g of (e) and processed into tablets with a tableting machine to give 1000 tablets each containing 10 mg of (a).

industrial Applicability

5

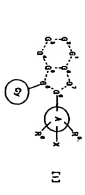
8 [0304] As is evident from the above-mentioned results, the compound of the prosent invention shows a high inhibitory activity against HCV polymerase.

[0305] Therefore, the compound of the present invention can provide a pharmaceutical agent effective for the prophylaxis or treatment of hopstills C, best on the anti-HCV offect afforded by the HCV polymerase inhibitory activity. When used concurrently with a different anti-HCV eight, such as interferon, and/or an anti-Hilliammatory agent and the like, it can provide a pharmaceutical agent more affective for the prophylaxis or treatment of hepatitis C. Its high inhibitory activity specific to HCV polymerase suggests the possibility of the compound being a pharmaceutical agent with slight also effects, which can be used satisfy for humans.

[0306] This application is based on patent application No. 359008/1999 filled in Japan, the contents of which are hereby incorporated by reference.

ક Claims

A therapeutic agent for hepatits C, which comprises a fused ring compound of the following formula (i) or a pharmaceutically acceptable sat thereof as an active ingredient:



Ġ

â

Ł

a broken line is a single bond or a double bond.

8					80	
	Q ⁷	G5, G6, G6 and G9	ଜୁ	ą.	Q ²	คั
	is C(·R7), an oxygon atom, a suffur atom, or a nitrogen atom optionally substituted by R8,	are each independently a carbon atom or a nitrogen atom,	is C(-R ⁴) or a ntrogen atom,	le C(-R ³) or a nitrogen atom,	is C(-R²) or a nitrogen atom,	is C(∙R¹) or a nitrogen atom,

wherein R1, R2, R3 and R4 are each independently,

hydrogen atom,

(2) Cr., a literocyl.
(3) corbooyl.
(4) cyreno.
(5) entrocyl.
(6) cyreno.
(6) Cr., a likyl optionally substituted by 1 to 3 substituent(s) selected from the following group A.
(7) Cr., a likyl optionally substituted Cr., a likyl cyrenologoup (Cr., a likyl cyrenologoup A.)
(8) entrocyl.
(9) to 5 substituent(s) selected from the following group B.
(9) to 6 substituent(s) selected from the following group B.
(9) to 6 substituent(s) selected from the following group B.
(10) coopers, (Crt), Coo

EP 1 162 186 A1

```
ij
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5
                                                                                                                 55
                                                                                                                                                                                                                                                                                               8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ŧ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                R<sup>5</sup> and R<sup>6</sup> are each independently
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (1) hydrogen atom,
(2) helogen atom,
(3) cyeno,
(4) hitro,
(5) eryeno,
(6) mitho, C., a sikenoylamino,
(6) eryeno,
(7) optionally substituted C., a sikyla (as defined abovo),
(8) C., a sikenylamino C., a sikyla (as defined abovo),
(9) C., a sikenyl optionally substituted by 1 to 3 substituent(s) selected from the above group A,
(9) C.OCRes
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  wherein R<sup>ab</sup> is hydrogan atom or C<sub>1-8</sub> elfyl, (10) -CONH-(CH<sub>2</sub>)-re<sup>10</sup> wherein R<sup>ab</sup> is optionally substituted C<sub>1-8</sub> alfyl (as defined above), C<sub>1-8</sub> alkoxycarbonyl or C<sub>1-8</sub> alkanylamino and 1 is 0 or an integer of 1 to 8.

(11) -ORati

    hydrogen atom,
    hatogen atom,
    hatogen atom,
    optionally substituted C<sub>1-6</sub> alityl (as defined above) or
    Optionally

                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             wherein Res is hydrogen atom, C1-a sikyl or C4-14 sryl C1-4 sikyl, and
                                                                                                                                                                                  group D:
                                                                                                                                                                                                              wherein the heterocyclic group has 1 to 4 hetero-etom(s) selected from an oxygen atom, a nitrogen atom
and a sulfur atom,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     wherein Ratt is hydrogen atom or optionally substituted C<sub>1-8</sub> alkyl (as defined above)
(a) hydrogen atom,
(b) halogen atom,
(c) cyano,
(d) nitro,

    (6) heterocyclic group optionally substituted by 1 to 5 substituent(s) solected from the following group
D

    a group selected from the following group D,
    C<sub>B,14</sub> eryl optionally substituted by 1 to 5 substituent(e) selected from the following group D,
    C<sub>B</sub> oycloalityl optionally substituted by 1 to 5 substituent(e) selected from the following group D,
    C<sub>B,14</sub> aryl C<sub>1,6</sub> alkyl optionally substituted by 1 to 5 substituent(e) selected from the following group D,

    C<sub>B-14</sub> eryl.
    C<sub>D-6</sub> cryclosikyl or
    heterocryclic group (as defined above).

                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     wherein ring B is
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       each Z is independently
```

ring A is

wherein u and v are each independently an integer of 1 to 3,

(1) C₆₋₁₄ eryl.
(2) C₂₄ cyclosikyl.
(3) C₂₄ cyclosikenyl or
(3) C₂₄ cyclosikenyl or
(4) heterocyclic group having 1 to 4 heterostom(s) selected from an oxygen stom, a nitrogen stom and s

8	
	(10°) C_{20} cycloalkyl C_{10} alkyl optionally substituted by 1 to 6 substituent(s) selected from the above group B,
8	(9") hotorocycle C ₁₋₄ ally/ optionally substituted by 1 to 6 substituent(s) selected from the above group B. (9") C ₃₋₆ cyclosity/ optionally substituted by 1 to 6 substituent(s) selected from the above group B, or
	group B, (7") heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B,
Ĝ	(2") optionally substituted C ₁₋₄ alkyl (as defined above), (3") optionally substituted C ₁₋₄ alkyl (as defined above), (4") C ₂₋₄ alkylyl optionally substituted by 1 to 3 substituent(s) selected from the above group A, (4") C ₂₋₄ alkylyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (5") C ₃₋₄ alyl optionally substituted by 1 to 5 substituent(s) selected from the above group B,
8	(1') hydrogen alom,
35	(I) -(CH ₂)-C(=NR=SD)NH ₂ Thorsion RSS is bettering a from or C = silvel
	b, or (87) Cy ₆ cycloally/ C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) solocied from the above group B.
8	wherein the heterocycle C ₁₋₈ alkyl is C ₁₋₈ alkyl substituted by heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group 8, as defined above, (77) C ₃₋₈ cycloally/ optionally substituted by 1 to 5 substituent(s) selected from the above group
	group B, (P) hotorocycle $C_{1:0}$ alkyl optionally substituted by 1 to 5 substituent(s) solected from the above only B.
	group B, (5") heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above
ર	(17) hydrogen stom, (27) optionally substituted C _{1,6} sityl (as defined above), (27) Optionally substituted by 1 to 5 substituent(s) selected from the above group B, (37) C ₆₋₁ , anyl C _{1,6} sityl optionally substituted by 1 to 5 substituont(s) selected from the above
	(h) -(CH ₂) ₁ -CONRe ²⁷ Re ²⁹ wherein Re ²⁷ and Re ²⁸ are each independently,
ដ	(g) -(CH ₂),-COORs ¹⁹ wherein Rs ¹⁸ is hydrogen storn, optionally substituted C ₁₋₆ sityl (as defined above) or C ₈₋₁₄ styl C ₁₋₆ sityl optionally substituted by 1 to 5 substituent(s) selected from the above group B.
70	group is wherein the heterocyclic group has 1 to 4 heterostom(s) selected from an oxygen stom, a nitrogen atom and a surfur stom,
	(1°) optionally substituted C _{1.4} alkyl (as defined above), (2°) C _{8.14} aryl optionally substituted by 1 to 5 substitutent(s) selected from the above group B or (3°) heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above
•	(o) optometry autostituted C1-g sitty! (as defined shows). (f) -(CH-b)-CO-Pers) (heroinafter each t means independently 0 or an integer of 1 to 6), wherein R118 is

EP 1 162 186 A1

8	:	80	å	à	Ł	30	t	8	8	ō	•		
substituted by 1 to 5 substituent(s) selected from the above group E, (8")-COORPA (RP3 is as defined above) or (7")-SO _A RPA (RP3 is as defined above),	or Jewen Heria hydrogen atom, optionally substituted C _{1,4} alty((as defined above), C _{8-1,4} ary) optionally substituted C _{1,4} alty((as defined above), C _{8-1,4} ary) optionally substituted by 1 to 6 autostituent(a) selocted from the above group B or C _{8-1,4} ary) C _{1,4} alty) optionally	(17) hydrogen atom, (27) optionally substituted C ₁₋₈ alkyl (as defined above), (37) C ₈₋₁₄ anyl C ₁₋₈ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (47) C ₈₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (57) CORSS	(11) O (OH ₂) ₀ , O (OH ₂) ₀	(e) -CO ₂ -(CH ₂) ₂ -NH-, (7) -CONH-(CH ₂) ₂ -NH-, (8) -NHCO ₂ -, (9) -NHCO ₂ -, (9) -NHCONH-,	(1) a single bond, (2) C _{1-a} altylane, (3) C _{1-b} altylane, (4) -(Ch ₂) _n -O-(CH ₂) _n . (4) -(Ch ₂) _n -O-(CH ₂) _n . (hereinfairer m and n are each independently 0 or an integer of 1 to 6), (5) -CO-,	w is an integer of 1 to 3, and Y is	(a). (CH ₂), SO ₂ , NHR ^{u28} (b). (CH ₂), SO ₂ , NHR ^{u28} (cH ₂), SO ₂ , SO ₂ (cH ₂), SO ₂ (cH ₂), SO ₂ , NHR ^{u28} (cH ₂), SO ₂ (cH ₂) (cH ₂), SO ₂ (cH ₂) (cH ₂), SO ₂ (cH ₂) (cH ₂	(n)-(CH ₂)-NHSO ₂ -R ⁴²⁵ wheels R ²² is hydrogen atom, optionally substituted C _{1-q} athyl (as defined above), C ₂₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B. (o)-(CH ₂)-S(O) _q -R ⁴²⁵ wherein R ⁴²⁵ is as defined above, and q is 0, 1 or 2.	(m) - (C+ ₂₋₃ ₁ -N;R ^{±28} CO-R ^{±24} wherein R ^{±28} is optionally substituted C₁ ₄ slityl or C₁ ₄ sittency, R ^{±24} is optionally substituted C₁ ₄ slityl (as defined above), C ₆₊₁ slyl optionally substituted by 1 to 5 substitutent(s) selected from the above group B or heterocyclic group optionally substituted by 1 to 5 substitutent(s) selected from the above group B.	group B or (5") heterocycle C_{14} sity/ optionally substituted by 1 to 5 substituent(e) selected from the above group B,	(1") hydrogen stom, (2") optionally substituted C ₁₋₀ shyl (as defined above), (3") C ₆₋₁₄ snyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, (4") C ₆₋₁₄ snyl C ₁₋₈ sityl optionally substituted by 1 to 5 substituent(s) selected from the above	() -(CH ₃),പ്യപ്പ്യമുള്ള wherein R ^{azz} and R ^{azz} as each independently	EFF I IVA I DO NI

(k) -(CH₂)₁-O- (CH₂)₂-COR^{Q1} wherein R^{Q2} is G_{1,0} altytemine or heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B, and p is 0 or an integer of 1 to 6,

(14) -NRe12CO- (Re12 is as defined above), (15) -CONRe13-(CH₂)_h-

nerein Re¹³ is hydrogon atom, optionally substituted C₁₋₄ alkyl (as defined above) or C₄₋₁₄ anyl C₁₋₄ alkyl ptionally substituted by 1 to 5 substituent(s) selected from the above group B.

arein R^{a 1}s is C₆₋₁₄ aryl optionally substituted by 1 to 5 substituent(e) selected from the above group 8, Ty -O-(CH₂)_m-CPA¹⁸FA¹⁸-(CH₂)_n-erein R^{a 18} and R^{a 18} are each independently

(1') hydrogen atom, (2') cerboxyl, (2') C-poxyl, (3') -Cpc (4') -Cpc (4') -Cpc (4') -Cpc (4') -Cpc (4') -Cpc (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P² (5') -NH2P²

wherein n', dng B', Z' and w' are the same as the above-mentioned n, dng B, Z and w, respectively, and may be the same as or different from the respective counterparts.

(18) -(CH₂)_A-NRe¹²-CHRe¹³- (Re¹³ and Re¹³ are each as defined above).
(19) -NRe¹⁷SO₂(19) -NRe¹⁷SO₂(20) -S(O)_e-(CH₂)_A-CR¹⁸SPe¹⁴-(CH₂)_A- (e is 0, 1 or 2, Re¹⁵ and Re¹⁶ are each as defined above).

- The therapoutic agent of claim 1, wherein 1 to 4 of the G1, G3, G3, G4, G5, G5, G7, G8 and G9 is (are) a nitrogen atom.
- The therapeutic agent of claim 2, wherein G^2 is $C(-R^2)$ and G^6 is a carbon atom.
- 4. The therapeutic agent of claim 2 or claim 3, wherein $\mathbf{G}^{\mathbf{5}}$ is a nitrogen atom.
- The thorapeutic agent of claim 1, wherein, in formuta (i), the moiety

is a fused ring selected from

EP 1 162 196 A1

The therapeutic agent of claim 5, wherein, in formula (i), the moisty

is a fused ring selected from

7. The therapeutic agent of claim 6, which comprises a fused ring compound of the following formula [1-1]

wherein each symbol is as defined in claim 1, or a pharmaceutically acceptable salt thereof as an active ingredient.

8. The therapeutic agent of claim 6, which comprises a fused ring compound of the following formula [I-2]

wherein each symbol is as defined in claim 1, or a pharmacoutically acceptable self thereof as an active ingredient.

The therapeutic agent of claim 6, which comprises a fused ring compound of the following formula [1-3]

EP 1 162 198 A1

wherein each symbol is as defined in claim 1, or a pharmaceutically acceptable salt thereof as an active ingredient.

10. The therapsutic agent of claim 8, which comprises a fused ring compound of the following formula [14]

wherein each symbol is as defined in claim 1, or a pharmecoutically acceptable selt thereof as an edition ingredient.

The therapeuric agent of any of claims 1 to 10, wherein at least one of A¹, A², A³ and A⁴ is carboxyl. COORs¹,
 -CONR¤2R³ or -SO₂R³ wherein R⁵¹, R²², R³ and R¹³ are as defined in claim 1.

The therapeutic agent of any of claims 1 to 11, wherein the ring Cy is cyclopentyl, cyclohexyl, cycloheptyl or tel-rehydrothlopyranyl.

13. The therapeutic agent of any of claims 1 to 12, wherein the ring A is Co-14 aryl.

14. A fused ring compound of the following formula [ii]

whorein the moiety

is a fused ring selected from

wherein R1, R2, R3 and R4 are each independently,

ŭ

(1) hydrogon atom,
(2) C₁₋₄ elkencyl,
(3) C₃-6 elkencyl,
(3) carboxyl,
(4) Cyano,
(5) nitro,
(6) nitro,
(6) C₁₋₄ elkyl optionally substituted by 1 to 3 substituent(s) selected from the following group A,
(7) -CODPa
(7) -CODPa
(7) -CODPa
(7) -CODPa
(7) -CODPa
(8) selected from the following group B,
(8) group B; halogon atom, oyano, nitro, C₁₋₄ alkyl, haloganated C₁₋₄ alkyl, C₁₋₄ alkyl, c₁₋₄ alkyl,
(7) -CODPa
(7) -(CH₂),-CODPa
(7) -(CH₂),-CODPa
(7) -(CH₂),-CODPa
(7) -(CH₂),-CODPa
(7) -(CH₂),-SO₂Pa

(8) -CONRP2R43 wherein R42 and R43 gro oach independently hydrogen atom, C₁₋₆ alkoxy or optionally substituted C₁₋₆ alkyl

(as defined above). (9) -C(=NRs4)NH₂ wherein Re4 is hydrogen atom or hydroxyl group.

Ġ

wherein \mathbf{R}^{ab} is hydrogen storn, \mathbf{C}_{1-b} alkanoyl or \mathbf{C}_{1-b} alkylaultonyl, (11) -OR ab

wherein R^{as} is hydrogen etom or optionally substituted C₁₋₆ alkyl (as defined above) . (12) -SO₂R^{a7} rherein $\bar{\mathsf{R}}^{\mathsf{a}\mathsf{T}}$ is hydroxyt group, amino, $\mathsf{C}_{\mathsf{1-g}}$ elkyt or $\mathsf{C}_{\mathsf{1-g}}$ alkytamino

(13) -P(=O)(OR 23)₂ wherein Re²¹ is hydrogen atom, optionally substituted C_{1-8} alkyl (as defined above) or C_{0+1} anyl C_{1-8} alkyl optionally substituted by 1 to 5 substitutent(s) selected from the above group B, and

R? is hydrogen atom or optionally substituted C₁₋₀ sityl (as defined above),

ring Cy is

(1) C_{3-8} cyclosikyl optionally substituted by 1 to 5 substituent(s) selected from the following group C_{\ast}

EP 1 162 198 A1

group C; hydroxy/ group, halogan atom, $C_{i,d}$ alkyl and $C_{i,d}$ alkoxy, or (2)

wherein ${f u}$ and ${f v}$ are each independently an integer of 1 to 3,

ü

5

mg A' is a group selected from a group consisting of phenyl, pyridyl, pyrazinyl, pyrimidinyl, pyridazinyl, cyclohaxyl, cyclohaxenyl, furyl and thianyl, H^{a} and H^{a} are each independently

8

(1) hydrogen atom,
 (2) hatogen atom,
 (3) optionally attrallitized C₁₋₆ alityl (as defined above) or
 (4) hydroxyl group

ring B ts

3

C₉₋₁₄ anyl.
 C₉₋₁₅ pysibilityl or
 head of the property of the proper

each Z is independently

ક

(1) a group selected from the following group D.

(2) C_{b-14} styl optionally substituted by 1 to 5 substituent(s) selected from the following group D.

(3) C_{b-2} cyclosity/ optionally substituted by 1 to 5 substituent(s) selected from the following group D.

(4) C_{b-14} styl C₁₋₆ sityl optionally substituted by 1 to 5 substituent(s) selected from the following group D or

(5) heterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the following group D wherein the heterocyclic group has 1 to 4 heteroetom(s) selected from an oxygen atom, a nitrogen atom and a sulfur atom,

å

ä

(a) hydrogen atom, (b) halogen atom, (c) cyano, (d) ntro,

å

(e) optionally substituted $C_{1,0}$ sity/ (as defined above), (f) -(CH₂), COR¹⁰, (hereinafter each I means independently 0 or an integer of 1 to 6),

wherein Ra19 is

8

(1) optionally substituted C_{1.9} alkyl (as defined above).
 (2) C₀₋₁₄ anyl optionally substituted by 1 to 5 substitutent(s) selected from the above group B or
 (3) hotorccyclic group optionally substituted by 1 to 5 substituent(s) solocted from the above

8

graps B wherein the heterocyclic group has 1 to 4 heterostom(s) selected from an oxygen stom, a nitrogen atom and a suffur atom,

(1') hydrogen atom. (2') optionally substituted C ₁₋₄ alkyl (as defined above), (3') C _{p-14} anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B. (4') C _{p-14} anyl C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B or (5') heterocycle C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B.	(k) -{CH ₂ },-C-(CH ₂) ₂ -C-CR ^{±21} wherein Re ^{±1} is C _{1,4} sitylamino or hotorocyclic group optionally substituted by 1 to 5 substituent (s) selected from the above group B, and p is 0 or an integer of 1 to 6, () -{CH ₂ },-NR ^{±2} 2R ^{±22} () -{CH ₂ },-NR ^{±2} 2R ^{±23} are each independently	(8) hotorocycle C ₁₋₈ sixyl optionally substituted by 1 to 5 substituent(s) selected from the above group 8. (8) C ₂₋₈ orchalkyl optionally substituted by 1 to 5 substituent(s) selected from the above group 8, or (10°) C ₂₋₈ orchalkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B.	(5) C ₀₋₁₄ any optionally substituted by 1 to 5 substituent(e) selected from the above group B, (8) C ₀₋₁₄ any C ₁₋₈ alkyl optionally substituted by 1 to 5 substituent(e) selected from the above group B. (7) heterocyclic group optionally substituted by 1 to 5 substituent(e) selected from the above	 hydrogen atom, optionally substituted C₁₋₆ alkyl (as defined above), optionally substituted C₂₋₆ alkyny (as defined above), optionally substituted C₂₋₆ alkynyl (as defined above), C₂₋₆ alkynyl optionally substituted by 1 to 3 substituent(e) solected from the above group 	(i) -(CH ₂),-C(=NR ⁴²³)NH ₂ wherein ন ⁴²³ is hydrogen etom or C ₁₋₆ elky!, (i) -(CH ₂),-OR ⁴²⁰ whorein নি ⁴²⁰ is	wherein the heterocycle C ₁₋₄ alkyl is C ₁₋₄ alkyl substituted by heterocyclic group optionally substituted by 1 to 5 substitutently) selected from the above group B, as defined above, (7) C ₃₋₆ cycloalkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B, or (8) C ₃₋₆ cycloalkyl C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B,	(5') heterocyclic group optionally substituted by 1 to 5 substituent(e) selected from the above group B, (6') heterocycle C ₁₋₆ alkyl optionally substituted by 1 to 5 substituent(s) selected from the above group B.	 hydrogen storn, optionally substituted C₁₋₆ sity/ (as defined above), optionally substituted C₁₋₆ sity/ (as defined above), O₉₋₁₄ ary/ optionally substituted by 1 to 5 substituent(a) selected from the above group B, O₉₋₁₄ ary/ C₁₋₆ sity/ optionally substituted by 1 to 5 substituent(a) selected from the above group B. 	(g) -{CH ₂ } ₁ -COORe ¹⁸ wherein Re ¹⁸ is hydrogen atom, optionally substituted C ₁₋₄ alityl (as defined above) or C ₆₋₁₄ aryl C ₁₋₄ alityl optionally substituted by 1 to 5 substitutent(s) selected from the above group B, (h) -{CD4} ₁ -COMR ² Pe ²⁸ (h) -{CD4} ₂ -COMR ² Pe ²⁸ are each independently.	EP 1 162 196 A1
8	\$6 &	ŧ	æ	æ	28	8	æ	6	·	

ä

8

12

8

ü

õ

EP 1 162 198 A1

(m) -(CH₂)-NRx2CCO-Rx24 wherein Rx2 is hydrogon atom, C₁₋₄ alkyl or C₁₋₄ alkanoyi, Rx24 is optionally substituted C₁₋₄ alkyl (as defined above), C₂₋₁₄ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B or haterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B.

(n)-(CH₂)-NHSO₂-Rx23 wherein Rx23 is hydrogen atom, optionally substituted C₁₋₄ alkyl (as defined above), C₂₋₁₄ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B or haterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B, (o)-(CH₂)-S(O)₂-Rx25 wherein Rx25 is as defined above, and q is 0, 1 or 2, (1) a single bond,
(2) C₁₊ silviyene,
(3) C₂₊ alleviyene,
(4) (CH₂)_n-O(CH₂)_n,
(4) (CH₂)_n-O(CH₂)_n,
(5) CD₂, CH₂)_n-O(CH₂)_n,
(6) CD₂, CH₂)_n,
(7) CONH-(CH₂)_n,NH-,
(8) AHCO₂-,
(9) AHCO₂-,
(9) AHCO₂-,
(10) O-(CH₂)_n-O,
(11) O-(CH₂)_n-O,
(12) SD₂-,
(13) (CH₂)_n-P,
(12) SD₂-,
(13) (CH₂)_n-P,
(13) (CH₂)_n-P,
(14) CD-(CH₂)_n-P,
(15) (CH₂)_n-P,
(16) CD-(CH₂)_n-P,
(17) CD-(CH₂)_n-P,
(18) CD-(CH₂)_n-P,
(19) CD-(CH w is an integer of 1 to 3, and y is (1) hydrogen atom,
(2) optionally substituted C₁₋₂ alityl (as defined above),
(3) C₂₊₁ anyl C₁₋₂ alityl optionally substituted by 1 to 5 substituent(s) selected from the above group B,
(4) C₂₊₁ anyl optionally substituted by 1 to 5 substituent(s) selected from the above group B,
(5) -CORPS
(5) -CORPS
wherein RPS is hydrogen atom, optionally substituted C₁₋₂ alityl (as defined above), C₂₊₁ aryl optionally substituted by 1 to 5 substituent(s) selected from the above group B or C₂₊₁ aryl C₁₋₂ alityl optionally substituted by 1 to 5 substituent(s) selected from the above group B,
(6) -COCPS (RPS is as defined above),
(7) -SO₂RPS (RPS is as defined above). (ρ) -(Ci+₂)₁-SO₂-NHR²³⁸ atom, optionally substituted C₁₋₈ sityl (as defined above), C₀₋₁₄ aryl wherein Pres is hydrogen atom, optionally substituted by 1 to 5 substituent(s) selected from the above group B or hoterocyclic group optionally substituted by 1 to 5 substituent(s) selected from the above group B. (14) -NR*12CO- (R*12 is as defined above),
(15) -CONR*13-(CH₂)_n.

wherein R*12 is hydrogen atom, optionally substituted C_{1,4} sity/ (as defined above) or C_{9,14} ary/
C_{1,4} sity/ optionally substituted by 1 to 5 substituent(s) solocated from the above group 8,
(16) -CONH-CHR*14.

wherein R*14 is C₉₊₁₄ ary/ optionally substituted by 1 to 5 substituent(s) selected from the above group B, (17) -O- (CH₂)_m-CRa15Ra16-(CH₂)_n-

E

8

å

å

vhorein Rais and Rais are each independently

(1') hydrogen atom.
 (2') carboxyl.
 (3') C₁₋₆ alkyl.
 (4') -ORbe

wherein R^{tos} is C₁₋₈ alityl or C₆₋₁₄ aryl C₁₋₈ alityl, or (5) -NHRP7 wherein R³⁷ is hydrogen atom, C₁₋₈ alityl, C₁₋₈ alkanoyl or C₆₋₁₄ aryl C₁₋₈ alityloxycarbonyl, or R³¹⁸ is optionally (6)

$$-(OH_2)$$
 $=$ $-(OH_2)$ $=$ $-(OH_2)$ $=$ $-(OH_2)$

wherein n', ring B', Z' and w' are the same as the sbove-mentioned n, ring B, Z and w, respectively, and may be the same as or different from the respective counterparts.

(16) -(CH₂), NR=12-CHR=13. (R=12 and R=15 are each as defined above), (19) -NR=17SO $_2$ -

wherein R^{a17} is hydrogen atom or C_{1,6} albyl or (20) -S(O)₆-(CH₂)₆-CR^{a15}R^{a16}-(CH₂)₆-(e is 0, 1 or 2, R^{a15} and R^{a16} are each as defined above), or a pharmaceutically acceptable salt thereof.

18. The fused ring compound of claim 14, which is represented by the following formula [II-1]

wherein each symbol is as defined in claim 14, or a pharmaceutically acceptable sall thereof.

16. The fused ring compound of claim 14, which is represented by the following formula [II-2]

wherein each symbol is as defined in claim 14,

EP 1 162 196 A1

or a pharmaceutically acceptable salt theraof.

17. The fused ring compound of claim 14, which is represented by the following formula [II-3]

wherein each symbol is as defined in cialm 14, or a pharmacoutically acceptable salt thereof.

18. The tused ring compound of claim 14, which is represented by the following formula [II-4]

ŝ

wherein each symbol is as dofined in claim 14, or a pharmaceutically acceptable salt thereof.

The fused ring compound of any of claims 14 to 18, wherein at least one of R1, R2, R3 and R4 is carboxyl, -COOR41
or -SO₂R42 wherein R41 and R42 are as defined in claim 14, or a pharmaceutically acceptable salt thereof.

20. The fused ring compound of claim 19, wherein at least one of R1, R2, R2 and R4 is carboxyl or COORs1 wherein Rs1 is as defined in claim 14, or a pharmaceurically acceptable sait thereof.

The fused ring compound of claim 20, wherein R² is carboxyl and R¹, R³ and R⁴ are hydrogen atoms, or a pharmacourically acceptable salt thereof.

22. The fused ring compound of any of claims 14 to 21, wherein the ring Cy' is cyclopentyl, cyclohoxyl, cycloheptyl or letrahydrothopyranyl, or a pharmaceutically acceptable self thereof.

The fused ring compound of claim 22, wherein the ring Cy is cyclopentyl, cyclohexyl or cycloheptyl, or a pharma-cautically acceptable salt thereof.

24. The fused ring compound of any of claims 14 to 23, wherein the ring A' is phonyl, pyridyl, pyrazinyl, pyrimidinyl or pyridazinyl, or a pharmacoutically acceptable salt thereof.

25. The fused ring compound of claim 24, wherein the ring A' is phenyl or pyridyl, or a phermacautically acceptable

26. The fused ring compound of claim 26, wherein the ring A' is phenyl, or a pharmaceutically acceptable salt thereof

- 27. The lused ring compound of any of claims 14 to 26, wherein the Y is -(CH₂)_m-O-(CH₂)_n^{*}, -NHCO₂*, -CONIH-CHR^{a14}, -(CH₂)_m-NR^{a12}-(CH₂)_h-CONIH-CHR^{a14}, -(CH₂)_m-NR^{a12}-(CH₂)_h-CONIH-CHR^{a14}, -(CH₂)_h-Or-(CH₂)_h-NR^{a12}-CHR^{a14}, (wherein each symbol is as defined in claim 14), or a pharmaceutically acceptable salt thereof.
- The fused ring compound of claim 27, wherein the Y is (CH₂)_n-O-(CH₂)_n-O-(CH₂)_n-CR^{a15}Ra¹⁶-(CH₂)_n-(whorein each symbol is as defined in claim 14), or a pharmacoutically acceptable salt thereof.
- The fused ring compound of claim 28, wherein the Y is -(CH₂)_m-O-(CH₂)_b, wherein each symbol is as defined in claim 14, or a pharmacounically acceptable salt thereof.
- 30. The tused ring compound of any of claims 14 to 29, wherein the R2 is carboxyl, R1, R3 and R4 are hydrogen atoms the ring Cy is cyclopentyl, cyclohaxyl or cyclohoptyl, and the ring A1 is phonyl, or a pharmacourtically acceptable sat thereof.
- The fused ring compound of claim 14 or a pharmacourtically acceptable satt thereof, which is selected from the group consisting of

ettyl 2:[4-(3-tromophenoxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate,
244(3-tromophenoxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate,
ettyl 2:[4-(2-tromo-6-foliorobarzyloxy)phenyl]-2-cyclohexylbenzimidazole-5-carboxylate,
ettyl 2:[4-(2-d-mon-6-foliorobarzyloxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate,
2:[4-(2-d-horophenyl)-5-chlorobarzyloxy)phenyl]-1-cyclohexylbenzimidazole-5-carboxylate,
ettyl 2:[4-(2-d-horophenyl)-5-methoxybenzyloxy)phenyl]-1-cyclohaxylbenzimidazole-5-carboxylate,
ettyl 2:[4-(2-d-horophenyl)-5-methoxybenzyloxy)phenyl]-1-cyclohaxylbenzimidazole-5-carboxylate,
ettyl 2:[4-(2-d-horophenyl)-5-methoxybenzyloxylphenyl]-1-cyclohaxylbenzimidazole-5-carboxylate,
ettyl 1-cyclohexyl-2-(4-([6]-2-phenylhinyl]phenylbenzimidazole-5-carboxylate,
ettyl 1-cyclohexyl-2-(4-([6]-2-phenylhinyl]phenylbenzimidazole-5-carboxylic acid,
et-4-banzyloxyphenyl)-1-cyclopentyBenzimidazole-5-carboxylic acid,
et-4-banzyloxyphenyl)-1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-carboxylide acid,
ettyl 1-cyclopentyBenzimidazole-5-carboxemide,
ettyl 1-cyclopentyBenzimidazole-5-

1-cyclohexyl-2-(4-((4-fluorophenyl)-2-methyl-5-thiazolyl)-methoxylphenyllbenzimidazole-6-carboxylic ac-id, sthy 2-(4-thia(3-fluorophenyl)methoxyl-2-fluorophenyl)-1-cyclohexylbenzimidazole-6-carboxylate, 2-(4-thia(3-fluorophenyl)methoxyl-2-fluorophenyl)-1-cyclohoxylbenzimidazole-6-carboxylic acid, athyl 2-(4-bonzoylaminophenyl)-1-cyclohoxylbenzimidazole-5-carboxylate,

etry test criscopiant phrestocky (2 chaptopiant) pro-production (24 dears of the Control principle) of the Control principle of the Control princi

â

ety-contry oxyphony) - re-productive introduction, and production of the Act-background

2-(4-bonzyloxyphonyl)-1-cyclopontyl-5-nitrobenzimidazole, fi-sanino-2-(4-benzyloxyphonyl)-1-cyclopontyl-benzimidazole hydrochloride, fi-santylamino-2-(4-benzyloxyphonyl)-1-cyclopontyl-benzimidazole, fi-santylamino-2-(4-benzyloxyphonyl)-1-cyclopontyl-benzimidazole, fi-suttamoyl-2-(4-benzyloxyphonyl)-1-cyclopontyl-benzimidazole, fi-suttamoyl-2-(4-benzyloxyphonyl)-1-cyclopontyl-benzimidazole,

EP 1 162 186 A1

trans-4 (2-(4-benzyloxyphenyl)-6-carboxybenzimidazei-1-yijbyciohexan-1-oi,
trans-1-12-(4-benzyloxyphenyl)-6-carboxybenzimidazei-1-yij4-marboxyyciohexane,
2-(4-benzyloxyphenyl)-6-carboxyntalyt-1-cyclopenylbenzimidazois2-(4-oxyciohexylphenyl-2-arboxyntalyt-1-cyclopenylbenzimidazois-2-denboytic edid,
1-cyclopenyl-2-(4-(3,6-dichlorobenzyloxylphenyl|benzimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-(3),6-dichlorobenzyloxylphenyl|benzimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-(bhenylexabarnoylamino) phonyl| benzimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-(bhenylexabarnoylamino) phonyl| benzimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-diphenylexabarnoylamino) phonyl| benzimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl)-6-carboxybranimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl)-6-carboxybranimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl)-6-carboxybranimidazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl)-1-cyclopenyl-0-oxidiazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl)-1-cyclopenyl-0-oxidiazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl-6-carboxybranyl-1-cyclopenyl-0-oxidiazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl-1-cyclopenyl-0-oxidiazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl-1-cyclopenyl-0-oxidiazois-6-carboxytic edid,
1-cyclopenyl-2-(4-benzyloxyphenyl-1-cyclopenyl-0-oxidiazois-6-carboxytic edid, 1-cyclohoxyl-2-(4-(3,5-dichlorobenzylaxy)phenylibenzimidazole-6-carboxylic acid,
1-cyclohoxyl-2-(4-dighenylmethoxy)phenylibenzimidazole-6-carboxylic acid,
1-cyclohoxyl-2-(4-dighenylmethoxy)phenylibenzimidazole-6-carboxylic acid,
1-cyclohoxyl-2-(4-1-3,5-direct-bury)barzylohoxyl)benzimidazole-5-carboxylic acid,
1-cyclohoxyl-2-(4-1-2-diphenyl)-1-(4-mothyl-cyclohoxyl)benzimidazole-5-carboxylic acid,
1-cyclohoxyl-2-(4-1-2-diphenyl)-1-(4-mothyl-cyclohoxyl)benzimidazole-5-carboxylic acid,
1-cyclohoxyl-2-(4-1-qarbhivy)methoxylphenyllbenzimidazole-5-carboxylic acid,
1-cyclohoxyl-2-(4-1-diphenyl-1-cyclohoxylarbhidazole-5-carboxylic acid,
2-(4-2-biphenyl-2-1-diphenyl-1-cyclohoxylarbhidazole-5-carboxylic acid,
2-(4-barzylarphenyl-1-cyclohoxylarbhidazole-5-carboxylic acid,
1-cyclohoxyl-2-(4-dibenzylarbhoxylphenyl|benzimidazole-5-carboxylic acid, 2-(2-benzylozy-5-pyrlay)-1-cyclohexythenzimidazole-5-carboxylic acid.
1-cyclohexy/-2-24-(2-(3-4,5-timethoxythenyl)ethoxylphenyl)-benzimidazole-5-carboxylic acid.
2-(4-benzylozyhenyl)-1-(4,4-dimethybychohexyl)benzimidazole-5-carboxylic acid.
1-cyclohexyl-2-(4-12-(1-naphthylphoxy)phenzylibenzimidazole-5-carboxylic acid.
2-(4-(2-benzylozyphenoxy)phonyl)-1-cyclohexylibenzimidazole-5-carboxylic acid. 2-(4-benzoylmethoxyphenyf)-1-cyclohexylbenzimidazole-5-carboxylic acid,
1-cyclohexyf-2 (4-(3-diphenyfaropyloxy)phenyfigenzimidazole-5-carboxylic acid,
1-cyclohexyf-2-(4-(3-carboxylic acid,
1-cyclohexyf-2-(4-(2-phency)enzyloxy)phenyfi-1-cyclohexyficarimidazole-5-carboxylic acid,
1-cyclohexyf-2-(4-(2-phency)erioxy)phenyfibenzimidazole-5-carboxylic acid,
1-cyclohexyf-2-(4-(2-phenyficopyloxy)phenyfibenzimidazole-5-carboxylic acid, 2-(4-(N-bonzenesulfonyi-N-mathylamino)phenyli-1-cyclopentyi-benzimidazole-5-carboxylic acid, 2-(4-(N-benzyi-N-mathylamino)phenyli-1-cyclopentyibonzimidazole-5-carboxylic acid, 1-cyclohexyi-2-(4-phenethylphenyl)benzimidazole-5-carboxylic acid, 2-(4-((4-chloropheny))carbony/aminojphenyi)-1-cyclopenyi-benzimidazole-5-carboxylic acid, 2-(4-((4-ten-buly/phony))carbony/aminojphenyi)-1-cyclopenyibenzimidazole-6-carboxylic acid, 2-(4-((4-ten-buly/phony))carbony/aminojphenyi)-1-cyclopenyibenzimidazole-6-carboxylic acid, 2-(4-((4-benzyloxypheny))carbony/aminojphenyi)-1-cyclopenyibenzimidazole-6-carboxylic acid 24-(2-chlorobonzyloxy)phenyl)-1-cyclopentylbenzimidazolo-5-carboxylic acid, 244-(3-chlorobenzyloxy)phenyl)-1-cyclopentylbenzimidazole-5-carboxylic acid, 2-(4-benzyloxyphenyl)-3-cyclopentylbenzimidazole-5-carboxylic acid, 2-(4-benzyloxyphenyl)-3-cyclopentylbenzimidazole-5-carboxylic acid, 1-cyclopenn/-2-(4-(4-mathytbonzyloxy)phenyljbenzimidazole-5-carboxylic acid, 1-cyclopennyl-2-(4-(3,5-dimethyl-4-isoxazolylimethoxylphenyl)-benzimidazole-5-carboxylic acid, 12-(4-benzyloxyphenyl)-1-cyclopennylbenzimidazol-5-yl-carbonylaminoscetic acid, 2-(4-(benzenesultonylamino)phenyl}-1-cyclopentyfbenzimidazolo-5-carboxyfic acid, 1-cyclopentyl-2-(4-(3,5-dichtorophenyfbarbonylamino)phenyl}-benzimidazole-5-carboxyfic acid, 2-[4-(4-carboxybenzylbxy)phenyl]-1-cyclopentylbenzimidazole-5-carboxylic acid, 2-[4-(4-chiorobenzylbxy)phenyl]-1-cyclopentylbenzimidazole-5-carboxylic acid, ı -cyclohexyl-2 (4-(2-hydroxyphenoxy)pheny|benzimidazole-5-carboxylic acid, 1-cyclohoxyl-2 (4-(3-hydroxyphanoxy)pheny|benzimidazole-5-carboxylic acid, 1-cyclohexyl-2 (4-(2-methoxyphenoxy)pheny|benzimidazole-5-carboxylic acid 2-[4-(4-tert-buty/benzy/bxy)/phenyl]-1-cyclopenty/benz/midazolo-5-cerboxylic acid -cyclohexyl-2-(4-(5-phonylpontyloxy)phonyljbenzimidezole-5-cerboxylic ecid, l-cyclopentyl-2-[4- (4-trilluoromethylbenzyloxy) phenyl]-benzimidazole-5-carboxylic acid, -cyclopentyl-2-[4-[4-pyridylmethoxy]phenyl]benzłmidazole-5-carboxylic acid hydrochlonde, -cyclopentyl-2-[4-(4-methoxybenzyloxy)phonyl]banzimidezole-5-carboxylic acid,

30

29

yra seru: 2,44 (3-chloro-8- (3,4 5-irtmethoxyphenyl) benzyloxyjphenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid, 2,44 (2-chlorenyl)phenyyl)phenyylynenyly	2-(4- (pia (4-fluoropreny)methoxy)-2-fluorophony)-1-cyclohexy/benzimidazole-5-carboxylic acid, 2-(4-(4-benzyloxyphenoxy)-2-chloropheny)-1-cyclohexy/benzimidazole-5-carboxylic acid, 2-(4-(4-benzyloxyphenoxy)-2-chloropheny)-1-cyclohexy/benzimidazole-5-carboxylic acid, 2-(4-(4-benzyloxyphenoxy)-2-chliturormethypheny)-1-cyclohexy/benzimidazole-5-carboxylic acid, 2-(4-(3-chloro-6-(2-chliturormethypheny)-1-cyclohexylonzimidazole-5-carboxylic acid, 2-(4-(2Ft)-2-amino-2-phenyle-toxylohexyl-1-cyclohexylohazimidazole-5-carboxylic acid, 2-(4-(2-bphenylyloxyphenyl)-1-cyclohexylohazimidazole-5-carboxylic acid, 2-(4-(2-bphenylyloxyphenyl)-1-cyclohexylohazimidazole-5-carboxylic acid, 2-(4-(2-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid, 2-(4-(3-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenyl)-1-cyclohexylbenzimidazole-5-carbox-ylic acid, 3-(4-(1-art-butoxycarbonyl-4-plperidy))methoxylphenoxyl-phenylp	2-(4-tibus(-4-methypheny)/methoxy)phenyl)-1-cyclchexy/benzimidazole-5-carboxylic acid, 2-(4-tibus(-4-fluoropheny)/methoxy)phenyl)-1-cyclchexy/benzimidazole-5-carboxylic acid, 1-cyclchexyl-4-methyxy-2-(4-(3-pheny)propoxy)phenyl)-benzimidazole-5-carboxylic acid, 1-cyclchexyl-4-nytroxy-2-(4-(3-pheny)propoxy)phenyl-benzimidazole-5-carboxylic acid, 1-cyclchexyl-4-nytroxy-2-(4-(3-pheny)propoxy)phenyl-benzimidazole-5-carboxylic acid, 1-cyclchexyl-4-nytroxy-2-(4-(3-pheny)propoxy)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 2-(4-(2-(2-benzy)coxypheny)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 2-(4-(2-carboxymethyloxyphenoxy)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 2-(4-(3-carboxymethyloxyphenoxy)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)pheny)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)pheny)phenyl-1-cyclchexybenzimidazole-5-carboxylic acid, 1-cyclchexyl-2-(2-methyl-2-(4-tifluoropheny)phenyl-1-cyclohaxybenzimidazole-5-carboxylic acid, 1-cyclchexyl-2-(2-methyl-2-(4-tifluoropheny)phenyl-1-cyclohaxybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)phenyl-1-cyclohaybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)phenyl)-1-cyclohaybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)phenyl)-1-cyclohaybenzimidazole-5-carboxylic acid, 2-(4-(3-chbro-8-(4-methypheny)phenyl)-1-cyclohaybenzimidazole-5-carboxylic acid,	1 - cyclohoxyi-2-(4-12-(4-trilluoromethylphenyl)benzyloxyl-phenyl)benzinidezole-6-carboxylic acid. 2-(4-t)big (4-chiropphenyl)methoxylphenyl)-1-cyclohoxybonzinidezole-6-carboxylic acid. 1-cyclohoxyi-2-4-42-(4-methoxyphenyl)-1-cyclohoxybonzinidezole-6-carboxylic acid. 1-cyclohoxyi-2-4-42-(3-methoxyphenyl)-benzinidezole-6-carboxylic acid. 1-cyclohoxyi-2-4-42-(3-methoxyphenyl)-benzinidezole-6-carboxylic acid. 1-cyclohoxyi-2-4-(2-cychentyl)-1-cyclohoxyphenylphen	1-cyclohexyl-2-{4-(3-mothoxyphenoxylphenyf)benzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-propoxyphenoxylphenyf)benzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-propoxyphenoxyphenyf)benzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-(3-methy-2-butenyfoxy)phenoxylphenyf)-benzimidazole-6-carboxylic acid, 1-cyclohexyl-2-{4-(2-(3-methy-2-butenyfoxy)phenoxylphenyf)-benzimidazole-6-carboxylic acid, 1-cyclohexyl-2-{4-(2-(2-lopenyfoxyphenoxy)phenyfipenzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-(2-lopenyfoxyphenoxy)phenyfipenzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-(3-leopenyfoxyphenoxy)phenyfipenzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-(3-leopenyfoxyphenoxy)phenyfipenzimidazole-5-carboxylic acid, 1-cyclohexyl-2-{4-(2-(10,11-dihydro-5)+-dibenzolo,fjazepin-5-yf)sihoxyfphenyfibenzimidazole-5-carboxylic acid,
--	--	---	--	---

25

8

ā

õ

g

8

EP 1 162 196 A1

B.	50	å .	ŧ	됞	8	25	20	õ	õ	o.
1-cyclohasyl-244-24-chlorophienyl)-3-nitrobenzyloxylphenyl)-benzimidazole-6-carboxylic acid. 1-cyclohasyl-24-19-(4-sintallydropyranyloxy)phonoxybhenyl)-benzimidazole-6-carboxylic acid, 1-cyclohasyl-24-19-(4-sintallydropyranyloxy)phonoxybhenyl)-benzimidazole-6-carboxylic acid, 1-cyclohasyl-24-19-(1-metmyl-4-piperfdy)lmethoxylphonoxyl-phonzimidazole-5-carboxylic acid, 1-cyclohasyl-24-19-(1-metmyl-4-piperfdy)lmethoxylphonoxyl-phonzimidazole-5-carboxylic acid, 2-14-19-(2-thorobenzyloxy)phonoxylphonyl)-1-cyclohasylbenzimidazole-6-carboxylic acid, 1-cyclohasyl-24-19-(3-19-yndy)phonoxylphonylphonzimidazole-6-carboxylic acid, 1-cyclohasyl-24-19-(3-19-yndy)phonoxylphonylphonzimidazole-6-carboxylic acid,	2-(4-f(2S)-1-(4-ecotylaminophenyl)-2-pyrrolidinylimthoxyl-phenyl)-1-cyclohexythenzimidazole-5-carboxylic acid, acid, (4-chlorophenyl)-2-metry/4-thlazolyimethoxylphenyl)-1-cyclohexythenzimidazole-5-carboxylic acid, (4-chlorophenyl)-2-metry/4-thlazolyimethoxylphenyl)-1-cyclohexythenzimidazole-5-carboxytic acid, (2-(4-f)sig/3-fluoropheny/jmethoxylphenyl)-1-cyclohexythenzimidazole-5-carboxytic acid, (2-(4-f)sig/3-fluoropheny/jmethoxylphenyl)-1-cyclohexythenzimidazole-5-carboxytic acid.	2-[4-[3-(4-chlorobenzyloxy)phenoxy]phenyl)-1-cyclohexylbenzimidezole-5-certoxylic edd, 1-cyclohexyl-2-[4-(4-fluorobenzyloxy)phenoxy]phenyl)-benzimidezole-5-certoxylic edd, 1-cyclohexyl-2-[4-([(25)-1-(4-nitrophonyl)-2-pyrrolidinyl)-methoxy]phenyl]benzimidezole-5-certoxylic edd, 1-cyclohexyl-2-[4-([(25)-1-phenyl-2-pyrrolidinyl]methoxy]phenyl]-benzimidezole-5-certoxylic edd hydrochlo- ride,	1-cyclohexyl-2-[4-[3-[(1-methanasulfonyl-4-pheridy])methoxy]-phenoxy[phenyl]benzimidazole-5-cerboxylic acid, acid,	2-(4-(1(25) - 1-benzulvzy) - pyrowny) promyty pod y pyrody pyrody (4((25) - 1-benzy) - 2-(4-(1(25) - 1-benzy) - 2-(4-(1(25) - 1-benzy) - 2-(4-(1(25) - 1-benzy) - 2-(4-(25) - 2-	2:44;2: 4-chorophanyy->nuoroomayaayjjananyy-1-cyconasyubaramazon-bcarboxyic acro, 2:44;3: eachory-6:44-chiorophanyjbanzyloxyjbhanyj1-1-cycohasybbanzimidazola-5-carboxyic acid, 2:44;3-carbamyy-6:44-chiorophanyjbanzyloxyjbhanyj1-1-cycohasybanzimidazola-5-carboxyic acid, 2:44;3-carbamyy-6:44-chiorophanyjbanzyloxyjbhanyj1-1-cycohasybanzimidazola-5-carboxyic acid, 1-cycohasyb-2:44;2-dimethycarbamoyimothoxyjbhanoxyjbhanyi1-banzimidazola-5-carboxyic acid, 1-cycohasyb-2:44;2-dippendincarbomyimothoxyjbhanoxyjbhanyi1-banzimidazola-5-carboxyic acid, 1-cycohasyb-2:44;2-dippendincarbomyimothoxybhanyibhanyi-banzimidazola-5-carboxyic acid, 1-cycohasyb-2:44;2-dippendincarbomyimothoxybhanyibha	2-(4-(2-(4-catroxypneryl-)-c-thoroen.zyloxypneryl-1-f-ycionexy.cent.mioazole-0-catroxylic scid. 2-(4-((2S)-1-benzyloxycarbonyl-2-pyrrolidinyl mothoxy phenyl]-1-cyclohexylbenzimidazole-5-carboxylic scid. id. 2-(2-chloro4-(2-(4-irfluoromethylphenyl)benzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylic scid. 1-cyclohexyl-2-(4-(3-(2-pyridylmethoxy)phenyl)-benzimidazole-5-carboxylic scid.	2-(4-(2-bromo-6-mathoxypurery ty	244(3-([1-seaty/4-pperidy)]methoxy)phonoxy[phonyl]-1-cylpionsybeaticalindazole-5-catboxylis acid, 24(4(3-([1-seaty/4-pperidy)]methoxy)phonoxy[phonyl]-1-cylcionsybeatimidazole-5-catboxylis acid, 24(4(3-([1-seaty/4-pperidy)]methoxy]phonoxy[phonyl]-1-cylcionsybeatimidazole-5-catboxylis acid, 1-cylcionsyl-2-(4(3-(2-propylryfoxy)phonoxy[phonyl]benzimidazole-5-catboxylis acid, 1-cylcionsyl-2-(4-(3-(2-propylryfoxy)phonoxy[phonyl]-bonzimidazole-5-catboxylis acid, 1-cylcionsyl-2-(4-(3-(2-pyrdylryfioxy)phonoxy[phonyl]-bonzimidazole-5-catboxylis acid, 1-cylcionsyl-2-(4-(3-2-yyldylryfioxy)phonoxy[phonyl]-bonzimidazole-5-catboxylis acid, 1-cylcionsyl-2-(3-pyrdylryfioxy)-1-cylcionsyl-2-(4-(3-2-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-2-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-(3-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-(3-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-(3-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-(3-yyldylryfioxy)-1-cylcionsyl-2-(4-(3-(3-yyldylryfioxy)-1-cylcionsyl-3-(4-(3-(3-(3-(3-(3-(3-(3-(3-(3-(3-(3-(3-(3-	2-(4-13-chloro-6-(3-chlorophony))benzyloxy)pheny)-1-cyclonexylonzimdazole-5-carboxylic acid, 2-(4-13-chloro-6-(3-pyrldy)benzyloxy)pheny)-1-cyclonexylbenzimdazole-5-carboxylic acid, 2-(4-13-chloro-6-(4-huoropheny))benzyloxy)pheny)-1-cyclohexylbenzimdazole-5-carboxylic acid, 2-(4-(4-benzyloxy)benzoy)-3-fluoropheny]-1-cyclohexylbenzimidazole-5-carboxylic acid, 2-(4-(4-benzyloxy)benzyloxy)-1-cyclohexylbenzimidazole-5-carboxylic acid, 2-(4-(2-brome-5-chlorobenzy)oxy)pheny]-1-cyclohexylbenzimidazole-5-carboxylic acid, 2-(4-(2-brome-5-chlorobenzy)oxy)pheny]-1-cyclohexylbenzimidazole-5-carboxylic acid, 3-(4-(2-brome-5-chlorobenzy)oxy)pheny]-1-cyclohexylbenzimidazole-5-carboxylic acid, 3-(4-(2-brome-5-chlorobenzy)pheny]-1-cyclohexylbenzimidazole-5-carboxylic acid, 3-(4-(2-brome-5-chlorobenzy)pheny]-1-cyclohe	2-[4-[((2S)-1-benzyl-2-pyrrolidiny]]mathoxy]phenyl]-1-cyclohexylbenzimidezole-5-carboxylic acid hydrochlo- ride, 1-[4-[-3-chloro-6-(4-methyllhlophenyl)benzyloxy]phenyl]-1-cyclohexylbenzimidezole-5-carboxylic acid, 2-[4-[3-chloro-6-(4-methyllhlophenyl)benzyloxy]phenyl]-1-cyclohexylbenzimidezole-5-carboxylic acid, 2-[4-[3-chloro-6-(2-thlenyl)benzyloxy]phenyl]-1-cyclohexylbenzimidezole-5-carboxylic acid, 2-[4-[3-chloro-6-(2-thlenyl)benzyloxy]phenyl]-1-cyclohexylbenzimidezole-5-carboxylic acid,

E

8

à

ylic acd. 1-cycbhasyl-2-{4-{14-(4-carboxyphanyl)-2-mathyl-5-thlazoly]-mathoxyjphanyljbenzimidazola-5-carboxylic acid hydrochlorda. 1-cycbhasyl-2-{2-fluoro-4-{4-fluoro-2-(3-fluorobanzoyl)-banzyloxyjphanyljbanzimidazola-5-carboxylic acid,
2-(4-benzyloxycyclohexyl)-1-cyclohexylbenzimidazole-6-carboxylic acid hydrochloride, 2-(2-(2-biphonylyloxymathyl)-5-chlonyl-1-cyclohexylbenzimidazole-6-carboxylic acid, 2-(2-(2-biphonylyloxymathyl)-5-chlonyl-1-cyclohexylbenzimidazole-6-carboxylic acid, 2-(2-(2-biphonylyloxymathyl-6-duryl)-1-cyclohexyl-6-zhlozoyyl-2-hydroxymathyl-6-thlazolyl]mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-(4-(4-diuorophenyl)-2-hydroxymathyl-6-thlazolyl]mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-(4-(4-diuorophenyl)-2-hydroxymathyl-6-thlazolyl]mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-(4-(4-diuorophenyl)-2-hydroxymathyl-6-thlazolyl]mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-(4-diuorophenyl)-2-hydroxymathyl-6-thlazolyl mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-duryl-6-thlazolyl mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-2-duryl-6-thlazolyl mothoxylphonyl benzimidazole-6-carbox-1-cyclohexyl-6-thlazolyl mothoxylphonyl benzimidazole-6-carbox-1-cyclohexylphonylphonyl mothoxylphonyl
2-14-2-(4-chlorophenyl)-5-sulfamoylbenzyloxyjphenyl)-1-cyclohexylbenzimidezole-5-carboxylic acid trifluor- oscatate
2-[4-[3-(tert-bury/suttamoyr)-6-(4-chlorophenyr)benzyloxy}-phenyr}-1-cyclohexylbenzimidszole-5-carboxylic acid.
2-(4-(-)-(-)-(-)-(-)-(-)-(-)-(-)-(-)-(-)-
mothyl 2-(4-(2-(4-chlorophenyl)-5-mothylcarbamoy/benzy/bxy)-phenyl)-1-cyclohexylbenzimidazole-5-carbox- viete
owyjeus; methyl 2-[4-[5-carboxy-2-(4-chlorophenyt)benzyloxy]phenyt)-1-cyclohexytbenzimidazole-5-carboxylate hy- drochloride.
ory e. (* * *******************************
chionde, chionde,
methyl 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylate, 2-(4-(2-(4-chlorophenyl)-5-methoxybenzyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid hydro-
1-cyclohexyl-2-[4-[4-(4-pyridy'meihoxy)-6-pyrimidinyloxy)phenyl)-benzimidazole-5-carboxylic acid, 2-[4-[3-chlorophenyl)-6-pyrimidinyloxy)phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid,
2-(4-(2-(4-chlorophenyl)-4-(6-tetrazolyl)benzyloxyjphenyl)-1-cyclohaxylbenzimidazole-5-carboxylio acid, 2-(4-(4-banzyloxy-6-pyrimidinyloxy)phenyl]-1-cyclohaxylbenzimidazole-5-carboxylio acid,
בייקיק-יארים ייטרים אין אייקיקיים אין הייטרים אין הייטרים אין אייקיים אייקיקיים אייקיקיים אייקיקיים אייקיקיים 2-{4/2-(3-chloropheny)-4-mathylanino-1,3,5-triaz in-8-yloxy phany)-1-cyclohaxylonzimidazote-6-carboxyl- ic acid trifluoroscetate,
- ((
2-(4-(4-chlorophenyl)-2-mothyl-5-pyrimidinyl)methoxy[phenyl]-1-cyclohexylbenzimidazole-5-carboxyllo add hydrochlorde,
2-(4-f2-(4-chlorophenyl)-5-dimethylcarbamoylbenzyloxy]phenyl)-1-cyclohaxylbenzimidazole-5-carboxylic ac- id,
1-cyclohexyl-2-44-{(4-4-dimethylcarbernoylphenyl)-2-methyl-5-thiazolyl methoxy phenyl]benzimidazole- 5-carboxylic acid,
2-(4-{2-(4-chlorophenyl)-5-ethoxycarbonylbenzyloxy]phenyl)-1-cyclohexylbenzimidezole-5-carboxylic acid, 1-cyclohexyl-2-(4-(3-trifluoromethylphenoxy)phenylbenzimidezole-5-carboxylic acid,
2-(4-(((2S)-1-(4-dimethycarbamoyiphanyl)-2-pymoldinyl)-methoxylphanyl)-1-cyclohaxylbonzimidezolo- 5-carboxylic acid,
2-(4/4-(4-chlorobenzyloxy)piperidinojphenyl)-1-cyclohexylbenzimidazole-5-cerboxylic acid. 2-(4/3-((2-chloro-4-cyridy))methoxylphenoxylphenyl)-1-cyclohexylbenzimidazole-5-cerboxylic acid.
2-(4-(3-(4-chlorobonzyloxy)ptporidino[phonyi]-1-cyclohaxylbenzimidazoie-5-carboxylia acid, 2-(4-(4-chlorobonzyloxy)ptporidino[phonyi]-1-cyclohaxylbenzimidazoie-5-carboxylia acid,
1-cyclohoxyi-2-(4-(3-(3,5-dichloropheny))phenoxy)phenyi)-benzimidazole-5-carboxylic acid, 2-(4-(1-(4-chlorobenzyi)-4-piperdyloxy)phenyi)-1-cyclohexy/benzimidazole-5-carboxylic acid,
1-cyclohexyl-214-[3-1(2-methyl-4-thiazoly)]methoxy)phenoxy)-phenyl[benzimidazole-5-carboxylic acid, 1-cyclohexyl-2-14-[3-1(2,4-dimethyl-5-thiazoly)]methoxy)phenoxy)-phenyl]benzimidazole-5-carboxylic acid,
id, 2-(4-11-(4-chlorobenzyl)-3-oioerdyloxylohenyl)-1-cyclohexyloenzimidazola-5-carboxylic edd.
boxylic acid, 2-{4-{\d-chlorophenyl) -2-methyl-5-thiazolyijmethoxyjphenyl}-1-cyclohexylbenzimidazole-5-carboxylic ac-
1-cyclohexyl-2-(4-[3-(4-methoxypheny/)phenoxyjphenyl)-benzimidszole-6-carboxylic acid, 1-cyclohexyl-2-(4-[(4-(4-methanesulfonylphenyl)-2-methyl-6-thlazolyl)methoxyjphenyljbenzimidazole-6-car-

EP 1 162 196 A1

2-(4-(3-(carboxymethy/carbernoyl)-8-(4-chlorophenyl)benzyloxy}-2-fluorophenyl)-1-cyclohexylbenzimida-	
5-carboxylic acid hydrochloride,	
ooxylia ada nyarachianana mbalinaca monyibanzyinyyi-2-fitoronbanyii-1-tyriobayibanyiniasyola. 2-(4-12-(4-chianabanyih-5-thianambalinacamonyibanzyinyi-2-fitoronbanyii-1-tyriobayibanyih-anyiniasyola.	•
2-(4-(2-(4-chlorophenyl)-5-morpholinocarbonylbenzyloxy)-2-fluorophonyl)-1-cyclohexylbenzimidazole-5-car-	•
dazole-5-carboxylic acid hydrochibride,	
2-(4-)2-(4-chlorophenyl)-5-(4-hydroxyptjeridino)-carbonylbenzyloxyl-2-iluorophonyli-1-cyclohexylbenzimi-	
2-(4-(2- (4-chlorophenyl)-5-(2-hydroxyethyl)carbemoylbenzyloxy}-2-fluorophenyl)-1-cyclohexylbenzimide- zolo-5-carboxyllo acid hydroxhorida	•
5-carboxylic acid hydrochloride,	
2-(4-(2-(4-chlorophenyl)-5-(1-pyrrolidinyl)carbonylbenzyloxy[-2-fluorophenyl]-1-cyclohexylbenzimidazoia-	
boxylic acid hydrochlorids	
2-(4-(2-(4-chlorophenyl)-5-piporidinocarbonylbonzyloxy]-2-fluorophenyl}-1-cyclohexylbenzimidazole-5-car-	
2-14-12-14-cnioropnenyi)-o-asopropyicaroamoyibenzyioxy -z-riuoropnenyi]-1-cyclonexyibenzimidazole-5-car- boxylic acid hydrochodda	-
boxylic scid hydrochloride,	
2-(4-(2-(4-chlorophenyl)-5-diethylcarbamoyibenzyloxy}-2-fluorophenyl)-1-cyclohexylbenzimidazole-5-car-	
ic soid hydrochloride,	
nydrochloride, 2.14.12.14phloride, and heree illorydeminopopydrydrydrhondt 1phloridensimi	~
2-(4-(2-(4-chlorophenyl)-5-dimethylaminobenzyloxy)phenyl)-1-cyclohexylbenzimidazolo-5-carboxylic acid di	
lato hydrochloride,	
methyl 2-44-5-cerboxy-2-(4-chlorophenyl)benzyloxyl-2-(luorophenyl)-1-cyclohexylbonzimidazole-5-cerboxy	
z-(+-]a-(+-c-more)nieny)-S-memenesationy benzywsy (prieny) (- 1-cyclonexy benzymaszore-S-caroxy) ic acro hydrochlaride.	-
2-44-(2-(4-chlorophenyl)-5-dimethylsuffamoylbenzyloxylphenyl)-1-cyclohexylbenzimidezolo-5-carboxylic ecto	
5-carboxy/lo acid.	
2-{4-{2-(4-chlorophenyl)-5-methoxybenzyloxy]-2-fluorophenyl}-1-(4-tetrahydrothlopyranyl)benzimidazole-	3
zimidazole-5-carboxylic scid,	
drochloride, 2-(4-(3-dimethylcarbamoyl-8-(4-dimothylcarbamoylphenyl)-benzylphenyl)-1-cyclohexylben	
2-[4-[3-dimethylcarbemoyl-6-(3-pyridyf)benzyloxy]phenyl]-1-cyclohexylbenzimidazolo-5-carboxylic acid diby	
5-carboxylic acid hydrochloride,	
2-{4-{3-dimethylcs/bamoyl-8-{4-methanosulfonylphonyl}-bonzyloxy]phenyl}-1-cyclohoxylbonzimidazole-	•
id hydrochloride.	
2-14-13-cg handy t-6-(4-chigrophony) benzyloxyt-2-fuorophonyl-1-cyclohazylbanzimidazola-5-pahonyl-a-	
entrijenje objektivalne i produktivalne i prod	
2444244410mnhappy1.54imathylrachampulhangday.1.3.fliographamil.1.mshhappilandaya.fl.ga	
د الطرة الإستان المراجعة المر	•
ic sea injureationes	
1-cyclohoxyl-2-(4-[3-dimethylcarbemoyl-6-(4-methylthlophenyl)-benzyloxyjphenyljbenzimidazole-5-carboxy	
boxylic acid hydrochleride,	
1-cyclohexyl-2-(4-[3-dimethylcarbamoy+8-(4-trifluoromethylphenyl)bonzyloxylphonyljbenzimidazolo-5-car-	•
hydrochioride,	
edo. 2-(4-)2-(4-chlorophenyi)-5-dimethylcarbamovlbenzyloxylphenyi]-1cyclohexylbenzimidazole-5-carboxylic acid	
درام خدام-cmoropneny)-o-mernoxybenzynoxy[pneny]-۱-(aterranydromopyrany)]bonzimidazole-5-carboxylic مراجع	
2-(4-(2-(4-carboxyphenyi)-3-pyridyi)methoxyjphenyi)-1 cyclohexylbenzimidezole-5-carboxylic acid,	۰
	•
2-(4-(3-carbamoyi-6-(4-chlorophenyi)benzyloxyjphenyi)-1 cyclohexylbenzimidazole-5-carboxylic acid hydro	
circired, 2-i4-12-i4-chlorophenyli-5-methoxy0enzyloxylphenyli-1-cyclohexylbanzimidazola-4-carboxylic acid	
1-cyclohexyl-2-(4-(3-carboxy-5-(4-pyridylmethoxy)phenoxylphenyl]benzimidezole-5-carboxylic acid dihydro	_
acid dihydrochloride,	
1-cyclohexyl-2-[4-[3-dimethylcarbarnoyl-5-[4-pyridylmothoxy]-phenoxy]phenyl]benzimidazole-5-carboxylic	
z-1+14-(+-cinorophenyl)-5-methorybenzyloxylphenyl)-1-cyclohexylbenzimidszole-5-eutrona scio, 2-14-12-(4-chlorophenyl)-5-methorybenzyloxylphenyl)-3-cyclohexylbenzimidszole-4-carboxylic scid.	
6.14.10.14. ahia-mahamili B. mathawahamilan dahamili da malahamili basadan la sadianta asid	

chloride,	
2-(4-(2-(4-chlorophenyt)-5-mothoxybenzylthio)phenyl)-1-cyclohoxylbonzimidazole-5-carboxylic acid hydro-	
drachloride,	
2-(4-(5-(4-chlorophenyl)-2-methoxybenzylsulfonyl]phenyl]-1-cyclohexylbenzlmidezole-5-cerboxylic ecid hy-	
drachloride,	
2-(4-(5-(4-chlorophenyi)-2-mothoxybenzyisulinyijphonyi)-1-cyclohexyibenzimidazois-5-carboxyiic acid hy-	
2-(4-(5-smino-2-(4-chlorophenyl)benzyloxy)phenyl)-1-cydohexylbenzlmidazole-5-carboxylic acid.	
boxylic acid,	
2-(4-(2-(4-carbamoyiphoriyi)-5-(dimethylcerbamoyi)benzyloxy)-phenyi)-1-cyclohaxylbanzimidazole-5-car-	
2-(4-(2-(4-cerboxyphenyi)-5-methoxybenzyloxy)phenyi)-1-cyclohexybenzimidezote-5-cerboxylic ecid,	
2-(4-16-carboxy-2-(4-chlorophonyl)benzyloxy}-2-fluorophenyl)-1-cyclohexylbenzimidazolo-6-carboxylic acid,	
zole-5-cerboxylate,	
sodium 24442-(4-chlorophenyi)-5-(dimethyticarbampyi)benzyloxy}-2-fluorophenyi)-1-cyclohexyibenzimida-	
zole-5-carboxylate.	
methyl 2-(4-(2-(4-chlorophenyl) -5- (dimethylcarbamoyl) benzyloxy}-2-fluorophenyl)-1-cyclohexylbenzimids-	
sodium 2-[4-[2-thlerryl-3-thlerrylmethoxy]-2-fluorophenyl)-1-cyclohoxylbenzimidazole-5-carboxylate,	
ā	
2-(4-fbis (4-dimethylcarbamoylphenyl)methoxyl-2-fluorophenyl)-1-cyclohexylbenzimidazole-5-carboxylic ac-	
2-(4-(bis(3-pyridyi)methoxy)-2-iluorophenyi)-1-cyclohexyibenzimidazole-5-carboxyiic acid.	
ride,	
2-(4-[2-(4-chlorophenyi)-5-cyanobenzyloxy]phenyi)-1-cyclohexyibenzimidazole-5-carboxyiic acid hydrochlo-	
2-(4-[2-(4-chlorophenyi)-5-mothylaulilnylbenzyloxy]phenyi)-1-cyclohexylbenzimidezolo-5-cerboxylic ecid.	
2-(4-(2-(4-chtorophenyt)-5-methytihiobenzyloxy)phenyt)-1-cyclohexylbenzimidazole-5-carboxylic acid.	
2-(4-(2-(3-carboxyphenyl)-5-chlorobenzyloxy phenyl)-1-cyclohexylbenzimidazole-5-carboxylic acid.	
hydrochloride,	
2-(4-(3-chioro-6-(4-methoxymethylphenyl)benzyloxylphenyl)-1-cyclohexytbenzimidazole-5-carboxylic acid	
drochloride,	
2-(4-(3-chloro-6-(4-hydroxymethyphenyf)benzyloxyjphenyf)-1-cyclohexylbenzimidazole-6-carboxylic acid hy-	
2-(4-(2-carboxyethy/)phenyl)-5-chlorobenzyloxylphenyl)-1-cyclohexylbenzimidezole-5-carboxylic ecid,	
zore-o-caroexylic add nyozochonde,	

õ

Ķ A pharmaceutical composition comprising a fused ring compound of any of claims 14 to 31, or a pharmaceutically acceptable sait thereof, and a pharmaceutically acceptable carrier.

8

8

2-(4-benzyloxyphenyl)-3-cyclohexylimidazo[1,2-djbyridina-7-carboxylic acid, and 2-(4-[2-(4-chlorophenyl)-5-methoxybenzyloxylphenyl)-3-cyclohexy-3H-imidazo[4,5-b]pyridina-8-carboxylic methy 2-(4-12-(4-chloropheny)-5-methoxybenzyloxylphenyl)-1-cyclohoxyl-1-H-indole-5-carboxylate, 2-(4-2)-(4-chloropheny)-5-methoxybenzyloxylphenyl)-1-cyclohoxyl-1-H-indole-5-carboxylate acid. 2-(4-benzyloxypheny))-1-cyclopentyl-1-H-indole-5-carboxylate acid, ethyl 2-(4-benzyloxypheny))-3-cyclohoxy-Imidazo(1,2-e)pyridine-7-carboxylate, Imidazo(1,2-e)pyridine-7-carboxylate.

8

8

Ĝ

dazole-5-carboxylic acid hydrochloride,

8

boxylic acid hydrochloride,

2-(4-{2-(4-chlorophenyl)-5-(4-pyridythothytzarbarnoy)benzytaxyj-2-fluorophenyl)-1-cyclohexytbenzimida-zole-5-carboxylia add dibydrochlorida, 2-(4-(2-(4-chlorophenyf)-5-(cyclohexylmethylcarbemcyf)benzyloxyf-2-fluorophenyf)-1-cyclohexylbenzimida-zole-5-carboxyfic acid hydrochloride,

2-(4-(2-(4-chiorophenyi)-5-(N-benzyi-N-methylcarbamoyi)-benzyioxy}-2-fluorophenyi)-1-cyclohexyibenzimi-

ㅂ

8

2-(4-jbis(4-carboxyphenyi)methoxy)-2-fluorophenyi)-1-cyclohexyibanzimidazole-5-carboxyib acid, 2-(4-jbis(4-carboxyibenzihoxy)-2-fluorophenyi)-1-cyclohexyibenzimidazole-6-carboxyib acid, mathyl 2-(4-(2-(4-chlorophenyi)-5-(methylcarbarroy))benzyibanzimidazole-mathyl 2-(4-(2-(4-chlorophenyi)-6-(methylcarbarroy))benzyibanzimidazole-mathyl 2-(4-(2-(4-chlorophenyi)-6-(methylcarbarroy))benzyibanzimidazole-mathyl

2-(4-{5-chloro-2-(4-pyridyl)benzyloxy|-2-fluorophenyl)-1-cyclohexylbenzimidazole-5-carboxylic ecid hydro-

2-{4-{2-(4-chlorophenyi)-5-{benzyicarbemoyi)benzyioxyj-2-fluorophenyi}-1-cyclohexyibenzimidazole-5-car-

Ğ

8

33. A hepatitis C virus polymerase inhibitor comprising a fused ring compound of any of claims 1 to 31, or a pharma-courtestly acceptable satt thereof, and a pharmacourtestly acceptable carrier.

EP 1 182 196 A1

- 34. An enti-hopalitis C virus agent comprising a fused ring compound of any of dalms 1 to 31, or a pharmaceutically acceptable earl thereof, and a pharmaceutically acceptable carrier.
- 35. A therapeutic agent for hepatitis C comprising a fused ring compound of any of claims 14 to 31, or a pharmaceutically acceptable sait thereof, and a pharmaceutically acceptable carrier.
- 36. A method for treating hepatitis C, which comprises administering an effective amount of a fused ring compound of the formula [i] of claim 1 or a pharmaceutically acceptable saft thereof.
- 37. A method for inhibiting hepatitis C virus polymenses, which comprises administering an effective amount of a fused ring compound of the formule [i] of claim 1 or a pharmacoutically acceptable salt thereof.

õ

- Use of a fused ring compound of the formula [I] of claim 1 or a pharmaceutically acceptable selt thereof for the
 production of a pharmaceutical agent for treating hepatitis C.
- Use of a fused ring compound of the formula [i] of claim 1 or a pharmaceutically ecceptable self thereof for the
 production of a hepatitis C virus polymorese inhibitor.
- 40. A phermaceutical composition for the treatment of hapatitis C, which comprises a fused ring compound of the formula [1] of ciaim 1 or a pharmaceutically acceptable salt thereof, and a pharmaceutically acceptable carrier.

ક

- 41. A pharmaceutical composition for inhibiting hopatitie C virus polymerase, which comprisos a fused ring compound of the formula [i] of claim 1 or a pharmacoutically acceptable salt thereof, and a pharmacoutically acceptable carrier.
- 25 A commorcial package comprising a pharmacourteal composition of claim 40 and a written matter associated thorowith, the written matter stating that the pharmacourteal composition can or should be used for treating hepatitis

43. A commorcial package comprising a pharmaceutical composition of claim 41 and a written matter associated therewith, the written matter stating that the pharmaceutical composition can or should be used for inhibiting hepatitis C virus polymerase.

8

t

INTERNATIONAL SEARCH REPORT

TMATTER 235/14, 235/30, 401/04, 401/10, 401/13, 401/14, 407/12, 405/04, 409/14, 2075411/04, 411/13, 417/13, 401/04, 407/04, MININ/407, 21, 433, 437, 4439, 484, 4709, MININ/4723, 486, 486, 406, 51, 3377,

man desperantian person (desidention priem fallemed by chalifeadon probab)

781-07 C070009/13, 1819/1, 1819/10, 1819/10, 1819/10, 1819/13, 1819/14, 1819/13, 1819/14,

ornale dra base countried during the international starch (surse of dra base und, where practicable, starch torns used)
CAPILIS . REGISTRY (GTN)

DOCUMENTS CONSIDERED TO BE RELEVANT EP, 507650, A1 (SYNTHELABO S.A.),

TO OCTOBER, 1992 (07.10.92),

A PT, 2674853, A C.A. 2064924, A

E NO, 2901291, A C.A. 201989, A

E NO, 1053459, A C.A. 2019853, A

E NU, 65573, A C.US, 52800300, A Chatton of document, with infraring, when appropriate, of the related passages (CO. 97)-(4237). A.I. (ELLT. LILLAY AUTH COMPANY).

11 December. 1997 (11.12.97).

12 C. 2327296. A. 6 AU, 9732329. A. 6 AU, 973229. A 1-35, 38-43 1-35, 38-43

Synthi magnite of died forestern:

A maintain is but y to previous of the sir which is not maintain the magnitude in the particles of the sir which is not published on a dier to harmstead filling the sir which may have don't no platch delarid.

The decrease which may have don'ts no platch delarid. xic, 97/2516, A1 (GRANG GROUT LPT.),
17 (17.5);
18 (17.7);
19 (17.5);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17.7);
18 (17. 1-35, 36-43

and determinants are listed in the continuation of Box C. | See permit theirly street.

In imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezine of size forement:

The imagezin

Dra- of the actual completion of the interactional cearch
20 February, 2001 (20.02.01) t published price to the international filing data but him wherity data exhaused Data of exciling of the international search report 06 March, 2001 (06.03.01)

Japanese Patent Office

329

EP 1 162 196 A1

Z
ğ
\$
귷
Ĕ
CO CD
æ
ξ
ñ
æ
-

This immunitional search report that not been established to respect of curats chains under Article 17(1)(1) for the following consoner	
---	--

The inventions of claims 36 and 37 fall under the category of methods for treatment of the human body by therapy.

 Claims Nos:
 because they relate to pure of the international application that do not comply with the prescribed extent that no manningful informational search can be extend on a specifically. Casina Nos.;
 because they are dependent claims and are not dashed in accordance with the second and third sentences of Bair 6.4(s).

As all required additional search fren wern timely paid by the applicant, this
claims.

As all searchable claims could be rescribed without either justifying as additioned in, with Authority did not to the payment of any additional fee.

As only scope of the required additional search these vers through paid by the applicant, this internst only those claims for which these were paid, specifically claims Mos;

No required additional search flow were timely paid by the applicant. Consequently, this international search report is married to the invention time measurement to the chains; it is covered by estima New.

Remark on Freien The additional search fiest were accompanied by the applicant's protest. No protest accompanied the payment of additional rearch thes.